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PAPAYA RINGSPOT VIRUS GENES

Abstract:

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(54) Title: **PAPAYA RINGSPOT VIRUS GENES**

(57) Abstract: The present invention relates to the isolation and identification of nucleic acid sequences encoding the coat protein of papaya ringspot virus in the Kapoho (KA), Keau (KE), Thailand (TH), Brazil (BR), Jamaica (JA), Mexico (ME), Venezuela (VE), and Oahu (OA) strains, and the uses thereof to impart viral resistance to papaya plants. The present invention also relates to nucleic acid constructs containing individual or multiple papaya ringspot virus coat protein-encoding nucleic acid sequences, and host cells and transgenic plants and seeds containing such constructs. The present invention is also directed to a method of using such constructs to impart to plants resistance to papaya ringspot virus.

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PAPAYA RINGSPOT VIRUS GENES

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FIELD OF THE INVENTION

The present invention relates to the isolation and purification of nucleic acid sequences encoding for papaya ringspot virus coat proteins, a method of conferring resistance to papaya ringspot virus by transforming plants with a construct containing one or more isolated viral coat protein nucleic acid sequences, and transgenic plants and seeds transformed with such multiple virus nucleic acid constructs.

BACKGROUND OF THE INVENTION

15 Papaya (*Carica papaya L.*) is an important fruit crop grown widely in tropical and subtropical lowland regions (Manshardt, "Papaya in Biotechnology of Perennial Fruit Crops," ed. Hammerschlag, 21:489-511, CAB Int., Wallingford, UK (1992)). Worldwide, Brazil, India, and Mexico are the largest producers of papaya. Hawaii, the largest producer of papaya in the United States, exports 66%
20 of the total fresh production, primarily to the U.S. mainland and to Japan (Martin, "Papaya Production Statistics," Proc. Annu. Hawaii Papaya Ind. Assoc. Conf., 39th, Kihei, pp. 31-36, Sept. 23-24 (1994)). In total production, papaya ranks above strawberries and below grapefruit (Manshardt, "Papaya in Biotechnology of Perennial Fruit Crops," ed. Hammerschlag, 21:489-511, CAB Int., Wallingford, UK (1992)). The FAO estimated that about 5.7 million metric tons of fruit were harvested in 1995, almost double the 1980 harvest (Galinsky, "World Market for Papaya," Reg. Agribus. Proj. Mark. Inf. Bull. Feb. No. 12, 5 pp. (1996)).

30 Papaya ringspot virus ("PRSV") is a member of the potyvirus group of plant viruses, which are pathogenic to several crop plants, and which exhibit cross-infectivity between members of different plant families. Generally, a potyvirus is a single-stranded (+) RNA plant virus. The viral genome is approximately 10,000 bases in length. The expression strategy of potyviruses includes translation of a complete polyprotein from the positive sense viral

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genomic RNA. PRSV is by far the most widespread and damaging virus that infects papaya, occurring worldwide wherever papaya is grown (Purcifull, "Papaya Ringspot Virus," CMI/AAB Descr. Plant Viruses, No. 292 (No. 84 Revis., July 1984) 8 pp. (1984)). PRSV infections have resulted in the devastation 5 of the papaya industry in Brazil, Taiwan, and Hawaii in recent years (Gonsalves, D., "Control of Papaya Ringspot Virus in Papaya: A Case Study," Annu. Rev. Phytopathol. 36:415-37 (1998)). Various attempts have been made to control or prevent infection of crops by PRSV, but these have been largely unsuccessful.

The concept of parasite-derived resistance ("PDR"), conceived in 10 the middle 1980s, offered a new approach for controlling PRSV (Sanford et al., "The Concept of Parasite-Derived Resistance - Deriving Resistance Genes from the Parasite's Own Genome," J. Theor. Biol. 113:395-405 (1985)). Parasite-derived resistance is a phenomenon whereby transgenic plants containing genes or sequences of a parasite are protected against detrimental effects of the same or 15 related pathogens. The application of PDR for plant viruses was first demonstrated when transgenic tobacco expressing the coat protein gene of tobacco mosaic virus was protected against infection by tobacco mosaic virus (Powell-Abel et al., "Delay of Disease Development in Transgenic Plants that Express the Tobacco Mosaic Virus Coat Protein Gene," Science, 232:738-43 (1986)). 20 Subsequent reports have shown that this approach is effective in controlling many plant viruses (Lomonossoff, G.P., "Pathogen-Derived Resistance to Plant Viruses," Ann. Rev. Phytopathol. 33:323-43 (1995)).

The vast majority of reports regarding PDR have utilized the coat 25 protein genes of the viruses that are targeted for control. Although the testing of transgenic plants have been largely confined to laboratory and greenhouse experiments, a growing number of reports have shown that resistance is effective under field conditions (Grumet, R., "Development of Virus Resistant Plants via Genetic Engineering," Plant Breeding Reviews 12:47-49 (1994)). Two virus resistant crops have been deregulated by the Animal and Plant Health Information 30 Service of the United States Department of Agriculture ("USDA/APHIS") and, thus, are approved for unrestricted release into the environment in the U.S. Squash that are resistant to watermelon mosaic virus 2 and zucchini yellow mosaic potyviruses have been commercialized (Fuchs et al., "Resistance of

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- Transgenic Hybrid Squash ZW-20 Expressing the Coat Protein Genes of Zucchini Yellow Mosaic Virus and Watermelon Mosaic Virus 2 to Mixed Infections by Both Potyviruses," Bio/Technology 13:1466-73 (1995); Tricoli, et al., "Field Evaluation of Transgenic Squash Containing Single or Multiple Virus Coat
- 5 Protein Gene Constructs for Resistance to Cucumber Mosaic Virus, Watermelon Mosaic Virus 2, and Zucchini Yellow Mosaic Virus," Bio/Technology 13:1458-65 (1995)). A transgenic Hawaiian papaya that is resistant to PRSV has also been developed (Fitch et al., "Virus Resistant Papaya Derived from Tissues Bombarded with the Coat Protein Gene of Papaya Ringspot Virus," Bio/Technology 10:1466-72 (1992); Tennant et al., "Differential Protection Against Papaya Ringspot Virus Isolates in Coat Protein Gene Transgenic Papaya and Classically Cross-Protected Papaya," Phytopathology 84:1359-66 (1994)). This resistant transgenic papaya was recently deregulated by USDA/APHIS. Deregulation of the transgenic papaya is timely, because Hawaii's papaya industry is being devastated by PRSV.
- 10
- 15 Remarkable progress has been made in developing virus resistant transgenic plants despite a poor understanding of the mechanisms involved in the various forms of pathogen-derived resistance (Lomonossoff, G.P., "Pathogen-Derived Resistance to Plant Viruses," Ann. Rev. Phytopathol. 33:323-43 (1995)). Although most reports deal with the use of coat protein genes to confer resistance,
- 20 a growing number of reports have shown that genes encoding viral replicase (Golemboski et al., "Plants Transformed with a Tobacco Mosaic Virus Nonstructural Gene Sequence are Resistant to the Virus," Proc. Natl. Acad. Sci. USA 87:6311-15 (1990)), movement protein (Beck et al., "Disruption of Virus Movement Confers Broad-Spectrum Resistance Against Systemic Infection by
- 25 Plant Viruses with a Triple Gene Block," Proc. Natl. Acad. Sci. USA 91:10310-14 (1994)), nuclear inclusion a-proteases ("Nla proteases") of potyviruses (Maiti et al., "Plants that Express a Potyvirus Proteinase Gene are Resistant to Virus Infection," Proc. Natl. Acad. Sci. USA 90:6110-14 (1993)), and other viral genes are also effective in conferring resistance. Furthermore, viral genes can be
- 30 effective in the translatable and non-translatable sense forms, and, less frequently, antisense forms (Baulcombe, D.C., "Mechanisms of Pathogen-Derived Resistance to Viruses in Transgenic Plants," Plant Cell 8:1833-44 (1996); Dougherty et al., "Transgenes and Gene Suppression: Telling us Something New?" Current

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Opinion in Cell Biology 7:399-05 (1995); Lomonosoff, G.P., "Pathogen-Derived Resistance to Plant Viruses," Ann. Rev. Phytopathol. 33:323-43 (1995)).

Notwithstanding the progress made in the field of plant resistance to viral pathogens, PRSV continues to exert its devastating effect upon papaya and other crops the world over. While the transgenic Hawaiian papaya is controlling the problem temporarily in Hawaii, that line unfortunately appears to susceptible to PRSV isolates with origins outside Hawaii. These observations suggest that transgenic papaya with coat protein genes specific to targeted PRSV isolates would need to be developed for transgenic papaya to effectively control PRSV worldwide. A more practical and comprehensive approach is needed to halt the devastation of PRSV. Such an approach would impart resistance to PRSV by utilizing genetic engineering techniques to provide greater and more reliable multi-pathogen resistance to crops to PRSV and other RNA-viral plant pathogens.

The present invention is directed to overcoming these and other deficiencies in the art.

SUMMARY OF THE INVENTION

The present invention relates to isolated nucleic acid molecules encoding a viral coat protein of papaya ringspot virus and the protein encoded by those nucleic acid molecules.

Another aspect of the present invention pertains to nucleic acid constructs containing the isolated nucleic acid molecules of the present invention operably linked to 5' and 3' regulatory regions.

The present invention also relates to nucleic acid constructs containing a plurality of trait DNA molecules, wherein at least some of the plurality of trait DNA molecules have a length that is insufficient to independently impart that trait to plants transformed with that trait DNA molecule. However, the plurality of trait DNA molecules are capable of collectively imparting their traits to plants transformed with the DNA construct and thereby effecting the silencing of the DNA construct. The trait associated with the DNA molecules of this construct is disease resistance, and the trait DNA molecules are derived from a gene encoding a papaya ringspot virus coat protein in a papaya ringspot virus

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strain selected from the group consisting of Thailand ("TH"), Keaau ("KE"), Kapoho ("KA"), Mexico ("ME"), Taiwan ("YK"), Brazil ("BR"), Jamaica ("JA"), Oahu ("OA"), and Panaewa ("PA").

The present invention also relates to a DNA construct containing a fusion gene which includes a trait DNA molecule which has a length insufficient to independently impart a desired trait to plants transformed with the trait molecule, operatively coupled to a silencer molecule effective to achieve post-transcriptional gene silencing. The trait DNA molecule and the silencer molecule collectively impart the trait to plants transformed with the construct. The DNA molecules of this DNA construct are derived from a gene encoding a papaya ringspot viral coat protein from a papaya ringspot virus strain selected from the group consisting of TH, KE, KA, ME, YK, BR, JA, OA, and VE.

The present invention also relates to host cells, plant cells, transgenic plants, and transgenic plant seeds containing the nucleic acid constructs of the present invention.

The present invention also relates to a method of imparting resistance against papaya ringspot virus to papaya plants. This involves transforming a papaya plant with the constructs of the present invention.

20

BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A-B show the cloning vectors used for the DNA constructs of the present invention. Figure 1A shows the expression cassette, pEPJ-YKT, containing the PRSV-CP variable regions of the YK, KE, and TH strains ligated into the pEPJ vector. Figure 1B shows the transformation vector pGA482G.

Figures 2A-B show the expression vectors used for cloning and subcloning the silencer-PRSV-CP construct. Figure 2A shows the pNP-YKT vector, containing the silencer DNA molecule (*M1/2NP*) and the *PRSV-CP* variable regions of PRSV strains YK, KE, and TH. Figure 2B shows the pGFP-YKT vector, containing the silencer molecule *GFP* ligated to the *PRSV-CP* variable regions of PRSV strains YK, KE, and TH PRSV strains.

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Figures 3A-G show various *PRSV-CP* DNA molecules ligated to the silencer molecule (*M 1/2 NP*) in an expression vector. Figure 3A shows clone pNP-K; Figure 3B shows clone pNP-KK; Figure 3C shows clone pNP-EE; Figure 3D shows clone pNP-KKTC; Figure 3E shows clone pNP-KKTV; Figure 3F shows clone pNP-EETC, and Figure 3G shows clone pNP-EETV.

Figure 4A shows the a full-length (1 Kb) *KE-CP* DNA molecule encoding a translatable RNA for PRSV-CP ligated into the expression vector pEPJ. Figure 4B shows a full-length (1 Kb) *KE-CP* DNA molecule encoding a non-translatable RNA for PRSV-CP ligated into the expression vector pEPJ.

10 Figure 5 shows a 855 bp *NcoI/BamHI* Mexico *PRSV-CP* DNA molecule ligated into the expression vector pEPJ.

DETAILED DESCRIPTION

15 The present invention relates to nucleic acids which encode for a viral coat protein ("CP") of papaya ringspot virus ("PRSV").

One suitable form of the nucleic acid of the present invention is the CP gene isolated from the PRSV strain Kapoho ("KA"), which has a nucleic acid sequence corresponding to SEQ ID NO: 1 as follows:

20 tccaaagaatg aagctgtgg a tgctggttt aatgaaaaac tcaaaggagaa agaaaagacag 60
aaagaaaaag aaaaagaaaa acaaaaagaa aaaggaaaag acgtatgtcg tgacgaaaat 120
gatgtgtcaa ctgcacaaaa aactggagag agagatagag atgtcaatgt tgggaccagt 180
ggaactttcg ctgttccgag aattaaatca ttactgata agttgattct accaagaatt 240
25 aaggaaaaga ctgtccttaa tttaagtcat cttcttcagt ataatccgca acaaattgac 300
atttctaaca ctcgtgccac tcagtcacaa tttgagaagt ggtatgaggg agtgaggat 360
gattatggcc ttaatgataa tgaatgcaa gttatgctaa atggtttgat gttttgggt 420
atcgagaatg gtacatctcc agacatatct ggtgtatggg ttatgatgga tggggaaacc 480
caagttgatt atccaaccaa gccttaatt gagcatgata ctccgtcatt taggcaaatt 540
30 atggctcact ttagtaacgc ggcagaagca tacattgcga agagaaatgc tactgagagg 600
tacatgccgc ggtacggaat caagagaaat ttgactgaca ttagcctcgc tagatatgct 660
ttcgacttct atgaggtgaa ttcaaaaaca cctgataggg ctcgcgaagc ccacatgcag 720
atgaaggctg cagcgctgctgaa acacactgt cgccagaatgt ttggatgga cggcagtgtt 780
agtaacaagg aagaaaacac ggagagacac acagtggaaat atgtcgatag agacatgcac 840
35 tctctcctgg gtatgcgcaaa ctat 864

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The present invention also relates to the PRSV-KA-CP, encoded by the nucleotide corresponding to SEQ ID NO: 1, where the protein encoded has an amino acid sequence corresponding to SEQ ID NO: 2, as follows:

5 Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Leu Lys Glu
1 5 10 15

Lys Glu Arg Gln Lys Glu Lys Glu Lys Glu Lys Gln Lys Glu Lys Gly
20 25 30

10 Lys Asp Asp Ala Ser Asp Glu Asn Asp Val Ser Thr Ser Thr Lys Thr
35 40 45

Gly Glu Arg Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Ala
15 50 55 60

Val Pro Arg Ile Lys Ser Phe Thr Asp Lys Leu Ile Leu Pro Arg Ile
65 70 75 80

20 Lys Gly Lys Thr Val Leu Asn Leu Ser His Leu Leu Gln Tyr Asn Pro
85 90 95

Gln Gln Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu
100 105 110

25 Lys Trp Tyr Glu Gly Val Arg Asp Asp Tyr Gly Leu Asn Asp Asn Glu
115 120 125

Met Gln Val Met Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly
30 130 135 140

Thr Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr
145 150 155 160

35 Gln Val Asp Tyr Pro Thr Lys Pro Leu Ile Glu His Asp Thr Pro Ser
165 170 175

Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile
180 185 190

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	Ala Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys			
	195	200	205	
5	Arg Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr			
	210	215	220	
10	Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln			
	225	230	235	240
	Met Lys Ala Ala Ala Leu Arg Asn Thr Ser Arg Arg Met Phe Gly Met			
	245	250	255	
15	Asp Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val			
	260	265	270	
	Glu Asp Val Asp Arg Asp Met His Ser Leu Leu Gly Met Arg Asn			
	275	280	285	

The present invention also relates to an isolated nucleic acid molecule encoding a *CP* gene isolated from the Thailand ("TH") strain of PRSV, which has a nucleic acid sequence corresponding to SEQ ID NO: 3 as follows:

	tccaagaatg aagctgtgga tgctggtctt aatgagaagt tcaaagataa agaaaaacag 60
25	aaagaagaaa aagataaaaca aaaaggtaaa gaaaataatg aagctagtga cgaaaaatgat 120
	gtgtcaacta gcacaaaaac tggagagaga gatagagatg tcaatgccgg aactagtgg 180
	actttcactg ttccgagaat aaaatttattt accgacaaga tgatttacc aagaattaag 240
	ggaaaaactg tccttagttt aaatcatctt cttcagtata atccgcaaca aatagacatc 300
	tcaaacactc gtgccactca atctcaattc gaaaagtgg atgaggagt gaggaatgat 360
30	tacggtctta atgataacga aatgcagaatg atgttaatg gtttgatggt ttgggtgcattc 420
	gaaaatggaa catccccaga catabctggt gtctgggtga tgatggatgg ggaaacccaa 480
	gtcgattatc ccatcaagcc tttgatcgaa catgcaactc cttcggttcag gcaaatcatg 540
	gctcaattca gtaacgcggc agaggcatac atcgcaaaaga ggaatgtac tgagaggtac 600
	atgcccgggt atggaatcaa gaggaatctg actgacattt gtctcgctag atatgctttc 660
35	gacttctatg aggtgaactc aaaaacaccc tataggggctc gtgaagctca tatgcagatg 720
	aaggctgcag cgctgcgcaa cactgatcgc agaatgtttg gaatggacgg cagtgtcagt 780
	aacaaggaag aaaacacccga gagacacaca gtgaaagatg tcaacagaga catgcactct 840
	ctccttaggtt tgcgcaattt a 861

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The present invention also relates to the viral coat protein of the TH strain of PRSV, encoded for by SEQ ID NO: 3, which corresponds to amino acid SEQ ID NO: 4, as follows:

5	Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Phe Lys Asp			
	1	5	10	15
	Lys Glu Lys Gln Lys Glu Glu Lys Asp Lys Gln Lys Gly Lys Glu Asn			
	20	25	30	
10	Asn Glu Ala Ser Asp Gly Asn Asp Val Ser Thr Ser Thr Lys Thr Gly			
	35	40	45	
	Glu Arg Asp Arg Asp Val Asn Ala Gly Thr Ser Gly Thr Phe Thr Val			
15	50	55	60	
	Pro Arg Ile Lys Leu Phe Thr Asp Lys Met Ile Leu Pro Arg Ile Lys			
	65	70	75	80
20	Gly Lys Thr Val Leu Ser Leu Asn His Leu Leu Gln Tyr Asn Pro Gln			
	85	90	95	
	Gln Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu Lys			
	100	105	110	
25	Trp Tyr Glu Gly Val Arg Asn Asp Tyr Gly Leu Asn Asp Asn Glu Met			
	115	120	125	
	Gln Val Met Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly Thr			
30	130	135	140	
	Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr Gln			
	145	150	155	160
35	Val Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser Phe			
	165	170	175	
	Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile Ala			
	180	185	190	

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	Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys Arg			
	195	200	205	
	Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr Glu			
5	210	215	220	
	Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln Met			
	225	230	235	240
10	Lys Ala Ala Ala Leu Arg Asn Thr Asp Arg Arg Met Phe Gly Met Asp			
	245	250	255	
	Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val Glu			
	260	265	270	
15	 			
	Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn			
	275	280	285	

Also suitable as a nucleic acid for use in the present invention is

20 the nucleic acid which encodes a *CP* gene isolated from the Keau ("KE") strain of PRSV. PRSV-KE contains two "cut-sites", i.e., two potential cleavage sites for a mature coat protein. The first cleavage site sequence in the KE strain of PRSV, identified herein as *KE-CP1*, corresponds to SEQ ID NO: 5 (*KECPI*) as follows:

	tcaaggagca ctgatgatta tcaacttgtt tggagtgaca atacacatgt gtttcattcag 60
	tccaaagaatg aagctgtgga tgctggttt aatgaaaaaac tcaaagagaa agaaaaacag 120
	aaaagaaaaag aaaaagaaaa acaaaaagaa aaaggaagag acgatgctag tgacgaaaat 180
	gatgtgtcaa ctgcacaaaa aactggagag agagatagag atgtcaatgt tgggaccagt 240
	gaaactttcg ctgttccgag aattaaatca ttactgata agttgattct accaagaatt 300
25	aaggaaaaga ctgtccttaa tttaagtcat cttcttcagt ataatccgca acaaattgac 360
	atttctaaca ctcgtgccac tcagtcacaa tttgagaagt ggttatgaggg agtgaggat 420
	gattatggcc ttaatgataa tgaaatgcaa gttatgctaa atggttgat ggtttggtgt 480
	atcgagaatg gtacatctcc agacatatct ggtgtatggg ttatgatgga tggggaaacc 540
	caagttgatt atccaaccaa gccttaatt gagcatgcta ctccgtcatt taggcaaatt 600
30	atggctcact ttagtaacgc ggcagaagca tacattgcga agagaaatgc tactgagagg 660
	tacatgccgc ggtacggaat caagagaaat ttgactgacg ttagcctcgc tagatatgct 720
	ttcgacttct atgaggtgaa ttcgaaaaca cctgataggg ctcgcgaagc ccacatgcag 780
	atgaaggctg cagcgctgca aaacactagt cgcagaatgt ttggatgga cggcagtgtt 840
	agtaacaagg aagaaaacac ggagagacac acagtggaaag atgtcaatag agacatgcac 900
35	tctctcctgg gcatgcgcaa c
	921
40	

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A second nucleotide sequence encoding a PRSV-KE coat protein sequence, which starts from the second KE-CP cleavage site, is identified as *KE-CP2* herein, and corresponds to SEQ ID NO: 6, as follows:

5

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tccaaagaatg aagctgtgga tgctggtttgc aatgaaaaac tcaaagagaa agaaaaacag 60
aaagaaaaaag aaaaagaaaa acaaaaaagaa aaaggaaaaag acgatgttag tgacgaaaat 120
gatgtgtcaa ctgcacaaa aactggagag agagatagag atgtcaatgt tgggaccagt 180
gaaactttcg ctgttccgag aattaaatca tttactgata agttgattct accaagaatt 240
10 aaggaaaaga ctgtccttaa tttaagtcat cttcttcagt ataatccgca acaaattgac 300
atttctaaca ctcgtgccac tcagtcacaa tttgagaagt ggtatgaggg agtgaggat 360
gattatggcc ttaatgataa tgaaatgcaa gttatgctaa atggtttgat ggtttgggt 420
atcgagaatg gtacatctcc agacatatct ggtgtatggg ttatgatgga tggggaaacc 480
caagttgatt atccaaccaa gccttaatt gagcatgcta ctccgtcatt taggcaaatt 540
15 atggctcaact ttagtaacgc ggcagaagca tacattgcgaa agagaaatgc tactgagagg 600
tacatgccgc ggtacggaat caagagaaat ttgactgacg ttagccctgc tagatatgct 660
ttcgacttct atgaggtgaa ttcgaaaaca cctgataggg ctcgcgaagc ccacatgcag 720
atgaaggctg cagcgctgca aaacactagt cgcagaatgt ttggatgga cggcagtgtt 780
agtaacaagg aagaaaaacac ggagagacac acagtggaaag atgtcaatag agacatgcac 840
20 tctctcctgg gcatgcgcaa cttaa 864
```

SEQ ID NOS: 5 and 6 contain, respectively, the N terminus and C terminus cleavage sites for PRSV-KE coat protein. Both cleavage sites result in proteins that appear to be functional in viral replication in the plant. SEQ ID NO: 5 encodes the first coat protein cleavage site product, CP1, of the KE strain of PRSV. KE-CP1 has an amino acid sequence corresponding to SEQ ID NO: 7, as follows:

	Ser Arg Ser Thr Asp Asp Tyr Gln Leu Val Trp Ser Asp Asn Thr His			
30	1	5	10	15
	Val Phe His Gln Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu			
	20 25 30			
35	Lys Leu Lys Glu Lys Glu Lys Gln Lys Glu Lys Glu Lys Gln			
	35 40 45			
	Lys Glu Lys Gly Arg Asp Asp Ala Ser Asp Glu Asn Asp Val Ser Thr			
	50 55 60			

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	Ser Thr Lys Thr Gly Glu Arg Asp Arg Asp Val Asn Val Gly Thr Ser			
65	70	75	80	
5	Gly Thr Phe Ala Val Pro Arg Ile Lys Ser Phe Thr Asp Lys Leu Ile			
	85	90	95	
	Leu Pro Arg Ile Lys Gly Lys Thr Val Leu Asn Leu Ser His Leu Leu			
	100	105	110	
10	Gln Tyr Asn Pro Gln Gln Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln			
	115	120	125	
	Ser Gln Phe Glu Lys Trp Tyr Glu Gly Val Arg Asp Asp Tyr Gly Leu			
15	130	135	140	
	Asn Asp Asn Glu Met Gln Val Met Leu Asn Gly Leu Met Val Trp Cys			
	145	150	155	160
20	Ile Glu Asn Gly Thr Ser Pro Asp Ile Ser Gly Val Trp Val Met Met			
	165	170	175	
	Asp Gly Glu Thr Gln Val Asp Tyr Pro Thr Lys Pro Leu Ile Glu His			
	180	185	190	
25	Ala Thr Pro Ser Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala			
	195	200	205	
	Glu Ala Tyr Ile Ala Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg			
30	210	215	220	
	Tyr Gly Ile Lys Arg Asn Leu Thr Asp Val Ser Leu Ala Arg Tyr Ala			
	225	230	235	240
35	Phe Asp Phe Tyr Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu			
	245	250	255	
	Ala His Met Gln Met Lys Ala Ala Leu Arg Asn Thr Ser Arg Arg			
	260	265	270	

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Met Phe Gly Met Asp Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu
275 280 285

Arg His Thr Val Glu Asp Val Asn Arg Asp Met His Ser Leu Leu Gly
5 290 295 300

Met Arg Asn
305

- 10 SEQ ID NO: 6 encodes the second coat protein cleavage site product, CP2, of the KE strain of PRSV. KE-CP2 has an amino acid sequence corresponding to SEQ ID NO: 8, as follows:

ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Leu Lys Glu
15 1 5 10 15

Lys Glu Lys Gln Lys Glu Lys Glu Lys Glu Lys Gln Lys Glu Lys Gly
20 25 30

20 Lys Asp Asp Ala Ser Asp Glu Asn Asp Val Ser Thr Ser Thr Lys Thr
35 40 45

Gly Glu Arg Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Ala
50 55 60

25 Val Pro Arg Ile Lys Ser Phe Thr Asp Lys Leu Ile Leu Pro Arg Ile
65 70 75 80

Lys Gly Lys Thr Val Leu Asn Leu Ser His Leu Leu Gln Tyr Asn Pro
30 85 90 95

Gln Gln Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu
100 105 110

35 Lys Trp Tyr Glu Gly Val Arg Asp Asp Tyr Gly Leu Asn Asp Asn Glu
115 120 125

Met Gln Val Met Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly
130 135 140

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	Thr Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr			
145	150	155	160	
5	Gln Val Asp Tyr Pro Thr Lys Pro Leu Ile Glu His Ala Thr Pro Ser			
	165	170	175	
	Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile			
	180	185	190	
10	Ala Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys			
	195	200	205	
	Arg Asn Leu Thr Asp Val Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr			
15	210	215	220	
	Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln			
	225	230	235	240
20	Met Lys Ala Ala Ala Leu Arg Asn Thr Ser Arg Arg Met Phe Gly Met			
	245	250	255	
	Asp Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val			
	260	265	270	
25	Glu Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn.			
	275	280	285	

Another nucleic acid suitable in the present invention is the *CP*
 30 gene isolated from the Taiwan ("YK") strain of PRSV, corresponding to SEQ ID
 NO: 9, as follows:

	tctaaaaatg aagctgtgga taccggctcg aatgagaagc tcaaagaaaa agaaaaagcag 60	
	aaagaaaaag aaaaagataa acaacaagat aaagacaatg atggagctag tgacggaaac 120	
35	gatgtgtcaa ctagcacaaa aactggagag agagataggg atgtcaatgc cgaaactagt 180	
	ggaaccttca ctgttccgag gataaagtca tttaactgata agatgatctt accaagaatt 240	
	aaggaaaaaa ctgtccctaa tttaaatcat cttcttcagt ataatccgaa acaagttgac 300	
	atctcaaaca ctcgcgccac tcaatctcaa tttgagaagt ggtatgaggg agtgagaaat 360	
	gattatggcc ttaatgataa cgaaatgcaa gtaatgttaa atggtttgat gggttgggtgt 420	
40	atcgaaaatg gtacatctcc agatatatct ggtgtctggg ttatgatgga tggggaaacc 480	

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caagtcgatt atccccattaa acctttgatt gaacacgcaa ctccttcatt taggcaaatc 540
 atggctcact tcaagtaacgc ggcagaggca tacatcgca agaggaatgc aactgagaag 600
 tacatgccgc ggtatggaat caagagaaat ttgactgaca ttagtctgc tagatatgct 660
 ttcgatttct atgaggtgaa ttcgaaaaca cctgataggg ctcgtgaagc tcataatgcag 720
 5 atgaaggctg cagcgctacg caatactaata cgcaaaaatgt ttggaatgga cgccagtgtc 780
 agtaacaagg aagaaaaaacac ggagagacac acagtggaag atgtcaacag agacatgcac 840
 tctctcctgg gtatgcgcaa ttga 864

SEQ ID NO: 9 encodes the CP of the YK strain of PRSV which has an amino acid
 10 sequence corresponding to SEQ ID NO: 10, as follows:

	Ser	Lys	Asn	Glu	Ala	Val	Asp	Thr	Gly	Leu	Asn	Glu	Lys	Leu	Lys	Glu
	1			5				10					15			
15	Lys	Glu	Lys	Gln	Lys	Glu	Lys	Glu	Lys	Asp	Lys	Gln	Gln	Asp	Lys	Asp
		20			25			30								
20	Asn	Asp	Gly	Ala	Ser	Asp	Gly	Asn	Asp	Val	Ser	Thr	Ser	Thr	Lys	Thr
		35			40								45			
25	Gly	Glu	Arg	Asp	Arg	Asp	Val	Asn	Ala	Gly	Thr	Ser	Gly	Thr	Phe	Thr
		50			55								60			
25	Val	Pro	Arg	Ile	Lys	Ser	Phe	Thr	Asp	Lys	Met	Ile	Leu	Pro	Arg	Ile
		65			70						75			80		
30	Lys	Gly	Lys	Thr	Val	Leu	Asn	Leu	Asn	His	Leu	Leu	Gln	Tyr	Asn	Pro
		85			90								95			
35	Lys	Gln	Val	Asp	Ile	Ser	Asn	Thr	Arg	Ala	Thr	Gln	Ser	Gln	Phe	Glu
		100			105								110			
35	Lys	Trp	Tyr	Glu	Gly	Val	Arg	Asn	Asp	Tyr	Gly	Leu	Asn	Asp	Asn	Glu
		115			120								125			
40	Met	Gln	Val	Met	Leu	Asn	Gly	Leu	Met	Val	Trp	Cys	Ile	Glu	Asn	Gly
		130			135								140			
40	Thr	Ser	Pro	Asp	Ile	Ser	Gly	Val	Trp	Val	Met	Met	Asp	Gly	Glu	Thr
		145			150								155			160

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	Gln Val Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser			
	165	170	175	
5	Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile			
	180	185	190	
	Ala Lys Arg Asn Ala Thr Glu Lys Tyr Met Pro Arg Tyr Gly Ile Lys			
	195	200	205	
10	Arg Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr			
	210	215	220	
	Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln			
15	225	230	235	240
	Met Lys Ala Ala Ala Leu Arg Asn Thr Asn Arg Lys Met Phe Gly Met			
	245	250	255	
20	Asp Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val			
	260	265	270	
	Glu Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn			
	275	280	285	
25	Another nucleic acid suitable in the present invention is the <i>CP</i> gene isolated from the Mexico ("ME") strain of PRSV, corresponding to SEQ ID NO: 11, as follows:			
30	tccaagaatg aagctgtgga tgctggtttgc aatgaaaaac tcaaagaaaaa agaaaaaacag 60 aaagaaaaaag aaaaacaaaaa agaaaaagaa aaagacaatg ctagtgacgg aaatgatgtg 120 tcgacttagca caaaaactgg agagaaagat agagatgtca atgtcggAAC tagtggAACT 180 ttcactgttc cgagaattaa atcatttact gataagatga ttctaccgag aattaaggga 240 aagactgtcc ttaatttaaa tcatcttctt cagtataatc cgcaacaaat tgatatttct 300			
35	aacactcggt ccactcagtc acaatttgag aaatggatgt agggagtgtg gaatgattat 360 ggctctgaatg ataatgaaat gcaagtgtatc ctgaatggct tgatggtttgc gtgtatcgag 420 aatggatcat ctccagacat atctgggttttgc tggatggggaa aattcaagttt 480 gactatccaa tcaagccctt aattgagcat gctaccccgat catttaggca gattatggct 540 cacttttagta acgcggcaga agcatatattt gcaaagagaa atgccactgtg gaggtacatg 600 40 ccgcggatgtt gatatcaagag aaatggactt gacatttagcc tcgcttaggtt ccgtttcgat 660			

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ttctatgagg ttaattcgaa aacacctgat agggctcgcg aagctcacat gcagatgaaa 720
gctgcagcgc tgcaaacac tagtcgcaga atgtttggta tggcggcag tgtagtaac 780
aaggaagaaa acacggaaag acacacagtg gaagatgtca atagagacat gcactctctc 840
ctgggtatgc gcaac 855

5

SEQ ID NO: 11 encodes the CP of the ME strain of PRSV which has an amino acid sequence corresponding to SEQ ID NO: 12, as follows:

10	Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Leu Lys Glu
	1 5 10 15
20	Lys Glu Lys Gln Lys Glu Lys Glu Lys Gln Lys Glu Lys Glu Lys Asp
	25 30
35	Asn Ala Ser Asp Gly Asn Asp Val Ser Thr Ser Thr Lys Thr Gly Glu
	40 45
50	Lys Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Thr Val Pro
	55 60
65	Arg Ile Lys Ser Phe Thr Asp Lys Met Ile Leu Pro Arg Ile Lys Gly
	75 80
85	Lys Thr Val Leu Asn Leu Asn His Leu Leu Gln Tyr Asn Pro Gln Gln
	90 95
100	Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu Lys Trp
	105 110
115	Tyr Glu Gly Val Arg Asn Asp Tyr Gly Leu Asn Asp Asn Glu Met Gln
	120 125
130	Val Met Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly Thr Ser
	135 140
150	Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Ile Gln Val
	155 160

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	Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser Phe Arg		
	165	170	175
5	Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile Ala Lys		
	180	185	190
	Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys Arg Asn		
	195	200	205
10	Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr Glu Val		
	210	215	220
	Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln Met Lys		
	225	230	235
	240		
15	 		
	Ala Ala Ala Leu Arg Asn Thr Ser Arg Arg Met Phe Gly Met Gly Gly		
	245	250	255
	Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val Glu Asp		
20	260	265	270
	Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn		
	275	280	285
25	Another nucleic acid suitable in the present invention is the <i>CP</i> gene isolated from the Brazil ("BR") strain of PRSV, corresponding to SEQ ID NO: 13, as follows:		
	tccaaaaatg aagctgtgga tgctggtttg aatgaaaagc gtaaaagaaca agagaaaacaa 60		
30	gaagaaaaag aagaaaaaca aaaaaagaaa gaaaaagacg atgctagtta cgaaaaacgat 120		
	gtgtcaacta gcacaagaac tggagagaga gacagagatg tcaatgttgg gaccagtgga 180		
	actttcaactg ttccgagaac aaaatcattt actgataaga tgattttacc tagaattaag 240		
	ggaaaaactg tccttaattt aaatcatctg attcagtata atccgaača aattgacatt 300		
	tctaacactc gtgctactca atcacaattt gagaagtgtt acgaggaggt gaggaatgtat 360		
35	tatggcccta atgataatga gatgcaaata gtgctaaatg gtttcatgtt ttgggtgtatc 420		
	gaaaacggta catctccaga catabctggt gtctgggtta tgatggatgg ggaaacccag 480		
	gttgactatc caatcaagcc tttaattgag catgctactc cgtcgttag gcaaattatg 540		
	gctcatttca gtaacgcggc agaagcatac attacaaaga gaaatgtac tgagaggtac 600		
	atgccgcggt atggatcaa gagaatttg actgacattha gtcttgtag atatgcttcc 660		
40	gatttctatg aggtgaattc gaaaacacct gatagggctc gcgaaagctca catgcagatg 720		

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aaagctgcag cgctgcgaaa cactaatcgc agaatgtttg gatatggacgg cagtgttagt 780
 aacaaggaag aaaacacgga gagacacaca gtggaaatgt tcaatagaga catgcactct 840
 ctccctgggta tgcgcaactg a 861

- 5 SEQ ID NO: 13 encodes the CP of the BR strain of PRSV which has an amino acid sequence corresponding to SEQ ID NO: 14, as follows:

	Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Arg Lys Glu		
1	5	10	15
10	Gln Glu Lys Gln Glu Glu Lys Glu Glu Lys Gln Lys Lys Lys Glu Lys		
	20	25	30
15	Asp Asp Ala Ser Tyr Gly Asn Asp Val Ser Thr Ser Thr Arg Thr Gly		
	35	40	45
	Glu Arg Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Thr Val		
	50	55	60
20	Pro Arg Thr, Lys Ser Phe Thr Asp Lys Met Ile Leu Pro Arg Ile Lys		
	65	70	75
	Gly Lys Thr Val Leu Asn Leu Asn His Leu Ile Gln Tyr Asn Pro Gln		
	85	90	95
25	Gln Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu Lys		
	100	105	110
30	Trp Tyr Glu Gly Val Arg Asn Asp Tyr Gly Leu Asn Asp Asn Glu Met		
	115	120	125
	Gln Ile Val Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly Thr		
	130	135	140
35	Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr Gln		
	145	150	155
	Val Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser Phe		
	165	170	175

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	Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile Thr		
	180	185	190
5	Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys Arg		
	195	200	205
	Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr Glu		
	210	215	220
10	val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln Met		
	225	230	235
	Lys Ala Ala Ala Leu Arg Asn Thr Asn Arg Arg Met Phe Gly Met Asp		
	245	250	255
15	Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val Glu		
	260	265	270
	Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn		
20	275	280	285

Another nucleic acid suitable in the present invention is a *CP* gene isolated from the Jamaica ("JA") strain of PRSV, corresponding to SEQ ID NO: 15, as follows:

25	tctaaaaatg aagctgtgga tgctggttta aatgaaaaagc tcaaagaaaa agaaaaacag 60 aaagataaaag aaaaagaaaa acaaaaagat aaagaaaaag gagatgctag tgacggaaat 120 gatggttcga ctagcacaaa aactggagag agagatagag atgtcaatgt tgggaccagt 180 ggaacttcca ctgttccgag aattaaatca ttcactgata agatggttct accaagaatt 240
30	aaggaaaaaa ctgtccctaa tttaaatcat ctttttcagt ataatccaca acaaattgac 300 atttctaaca ctcgtgccac tcagtcacaa tttgagaagt ggtacgaagg agtgaggagt 360 gattatggcc taaatgatag tgaaatgcaa gtgacgctaa atggcttgcgtt ggtttgggtgt 420 atcgagaatgt gtacatctcc agacatatct ggtgtctggg ttatgatgga tggggaaacc 480 caagttgatt atccaatcaa gcctttaatt gagcacgcta ccccatcatt taggcagatt 540
35	atggctcact tcagtaacgc ggcagaagca tacactgcaa agagaaatgc tactgagagg 600 tacatgccgc ggtatggaat caagagaaat ttgactgaca ttagtctcgc tagatacgct 660 ttcgatttct atgaggtgaa ttcaagaca cctgataggg ctcgtgaagc tcacatgcag 720 atgaaagctg cagcgcgtcg aaacactaat cgccagaatgt ttggatgga cggcagtgtt 780 agtaacaatg aagaaaacac ggagagacac acagtggaaat atgtcttat agacatgcac 840
40	tctctcctgc gtttgcgca ctga 864

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SEQ ID NO: 15 encodes the CP of the JA strain of PRSV which has an amino acid sequence corresponding to SEQ ID NO: 16, as follows:

	Ser	Lys	Asn	Glu	Ala	Val	Asp	Ala	Gly	Leu	Asn	Glu	Lys	Leu	Lys	Glu		
5	1			5					10				15					
		Lys	Glu	Lys	Gln	Lys	Asp	Lys	Glu	Lys	Glu	Lys	Gln	Lys	Asp	Lys	Glu	
							20			25				30				
10	Lys	Gly	Asp	Ala	Ser	Asp	Gly	Asn	Asp	Gly	Ser	Thr	Ser	Thr	Lys	Thr		
								35			40			45				
		Gly	Glu	Arg	Asp	Arg	Asp	Val	Asn	Val	Gly	Thr	Ser	Gly	Thr	Ser	Thr	
								50			55			60				
15																		
		Val	Pro	Arg	Ile	Lys	Ser	Phe	Thr	Asp	Lys	Met	Val	Leu	Pro	Arg	Ile	
								65			70			75			80	
		Lys	Gly	Lys	Thr	Val	Leu	Asn	Leu	Asn	His	Leu	Leu	Gln	Tyr	Asn	Pro	
20								85			90			95				
		Gln	Gln	Ile	Asp	Ile	Ser	Asn	Thr	Arg	Ala	Thr	Gln	Ser	Gln	Phe	Glu	
								100			105			110				
25	Lys	Trp	Tyr	Glu	Gly	Val	Arg	Ser	Asp	Tyr	Gly	Leu	Asn	Asp	Ser	Glu		
								115			120			125				
		Met	Gln	Val	Thr	Leu	Asn	Gly	Leu	Met	Val	Trp	Cys	Ile	Glu	Asn	Gly	
								130			135			140				
30																		
		Thr	Ser	Pro	Asp	Ile	Ser	Gly	Val	Trp	Val	Met	Met	Asp	Gly	Glu	Thr	
								145			150			155			160	
		Gln	Val	Asp	Tyr	Pro	Ile	Lys	Pro	Leu	Ile	Glu	His	Ala	Thr	Pro	Ser	
35									165			170			175			
		Phe	Arg	Gln	Ile	Met	Ala	His	Phe	Ser	Asn	Ala	Ala	Glu	Ala	Tyr	Thr	
								180			185			190				

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	Ala Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys		
	195	200	205
5	Arg Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr		
	210	215	220
	Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln		
	225	230	235
10	Met Lys Ala Ala Ala Leu Arg Asn Thr Asn Arg Arg Met Phe Gly Met		
	245	250	255
	Asp Gly Ser Val Ser Asn Asn Glu Glu Asn Thr Glu Arg His Thr Val		
	260	265	270
15	 		
	Glu Asp Val Tyr Ile Asp Met His Ser Leu Leu Arg Leu Arg Asn		
	275	280	285

Another nucleic acid suitable in the present invention is a *CP* gene
 20 isolated from the Oahu ("OA") strain of PRSV, corresponding to SEQ ID NO: 17,
 as follows:

	tccaagaatg aagctgtgga tgctggtttgc aatgaaaaat tcaaagagaa ggaaaaacag 60	
	aaagaaaaag aaaaagaaaa acaaaaagag aaagaaaaag atggtgctag tgacgaaaat 120	
25	gatgtgtcaa ctgcacaaa aactggagag agagatagag atgtcaatgt cgggaccagt 180	
	ggaactttca cagttccgag aattaaatca tttactgata agatgattct accgagaatt 240	
	aaggggaaagg ctgtccttaa tttaaatcat cttcttcagt acaatccgca acaaatcgac 300	
	atttctaaca ctcgtgccgc tcattcacaa tttgaaaagt ggtatgaggg agtgaggaat 360	
	gattatgccccc ttaatgataa tgaaatgcaat gtgatgctaa atggtttgat ggtttgggt 420	
30	atcgagaatg gtacatctcc agacatatct ggtgtctggg taatgatgga tggggaaacc 480	
	caagtcgatt atccaatcaa gcctttgatt gagcatgcta ctccgtcatt taggcaaatt 540	
	atggctcaact ttagtaacgc ggcagaagca tacattgcga agagaaaatgc tactgagagg 600	
	tacatgccgc ggtatgaaat caagagaaaat ttgactgaca ttagcctcgc tagatacgt 660	
	ttcgactttt atgaggtgaa ttcgaaaaca cctgatagag ctcgcgaagc tcacatgcag 720	
35	atgaaggctg cagcgctgcaaa acacaccagt cgccagaatgt ttggatgga cggcagtgtt 780	
	agtaacaagg aagaaaacac ggagagacac acagtggaaat atgtcaatag agacatgcac 840	
	tctctcctgg gtagcgc当地 ctaa 864	

SEQ ID NO: 17 encodes the CP of the OA strain of PRSV which has an amino
 40 acid sequence corresponding to SEQ ID NO: 18, as follows:

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	Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Phe Lys Glu		
1	5	10	15
5	Lys Glu Lys Gln Lys Glu Lys Glu Lys Glu Lys Gln Lys Glu Lys Glu		
	20	25	30
10	Lys Asp Gly Ala Ser Asp Glu Asn Asp Val Ser Thr Ser Thr Lys Thr		
	35	40	45
15	Gly Glu Arg Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Thr		
	50	55	60
20	Val Pro Arg Ile Lys Ser Phe Thr Asp Lys Met Ile Leu Pro Arg Ile		
	65	70	75
25	Lys Gly Lys Ala Val Leu Asn Leu Asn His Leu Leu Gln Tyr Asn Pro		
	85	90	95
30	Gln Gln Ile Asp Ile Ser Asn Thr Arg Ala Ala His Ser Gln Phe Glu		
	100	105	110
35	Lys Trp Tyr Glu Gly Val Arg Asn Asp Tyr Ala Leu Asn Asp Asn Glu		
	115	120	125
40	Met Gln Val Met Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly		
	130	135	140
45	Thr Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr		
	145	150	155
50	Gln Val Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser		
	165	170	175
55	Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile		
	180	185	190
60	Ala Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys		
	195	200	205

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	Arg Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr			
	210	215	220	
	Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln			
5	225	230	235	240
	Met Lys Ala Ala Ala Leu Arg Asn Thr Ser Arg Arg Met Phe Gly Met			
	245	250	255	
10	Asp Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val			
	260	265	270	
	Glu Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn			
	275	280	285	
15				

Another nucleic acid suitable in the present invention is the *CP* gene isolated from the Venezuela ("VE") strain of PRSV, corresponding to SEQ ID NO: 19, as follows:

20	atggctgtgg atgctggttt gaatggaaag ctcaaagaaa aagagaaaaa agaaaaagaa 60 aaagaaaaac agaaagagaa agagaaaagat gatgctagt acggaaatga tgtgtcaact 120 agcacaaaaa ctggagagag agatagagat gtcaaatattt ggaccagtgg aactttcact 180 gtcccttagga ttaaatcatt tactgataag atgattttac cgagaattaa gggaaagact 240 gtccttaatt taaatcatct tcttcagttt aatccgaaac aaatttgacat ttctaatact 300
25	cgtgccactc agtcgcatt tgagaaatgg tatgaggagat tgagggatga ttatggcatt 360 aatgataatg aaatgcattt gatgctaaat ggcttgatgg ttttgtcat tgagaatgg 420 acatctccag acatatctgg tttttgggtt atggtgatg gggaaaccca agttgattat 480 ccaatcaagg ctttaattga gcatgctaca ccgtcattt ggcaaattat ggctcatttt 540 agtaacgcgg cagaagcata cattgcgtt agaaatgcta ctgagaggta catgccgcgg 600
30	tatgaaatca agagaaattt gactgacatc aacctagctc gatacgctt tgatttctat 660 gaggtgaatt cggaaacmcc tggataggct cgtgaagctc acatgcagat gaaggctgca 720 gctttgcgaa acactaatcg cagaatgttt ggtatcgacg gcagtgttag caacaaggaa 780 ggaaacacgg agagacacac agtggatgtt gtcaatagag acatgcactc tctcctgggt 840 atgcgcactt aaatactcgc acttggatgtt ttgtcgagcc tgact 885
35	

SEQ ID NO: 19 encodes the CP of the VE strain of PRSV which has an amino acid sequence corresponding to SEQ ID NO: 20, as follows:

	Met Ala Val Asp Ala Gly Leu Asn Gly Lys Leu Lys Glu Lys Glu Lys			
40	1	5	10	15

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	Lys	Glu	Lys	Glu	Lys	Glu	Lys	Gln	Lys	Glu	Lys	Glu	Lys	Asp	Asp	Ala
	20							25						30		
5	Ser	Asp	Gly	Asn	Asp	Val	Ser	Thr	Ser	Thr	Lys	Thr	Gly	Glu	Arg	Asp
		35						40						45		
	Arg	Asp	Val	Asn	Ile	Thr	Ser	Gly	Thr	Phe	Thr	Val	Pro	Arg	Ile	Lys
		50						55				60				
10	Ser	Phe	Thr	Asp	Lys	Met	Ile	Leu	Pro	Arg	Ile	Lys	Gly	Lys	Thr	Val
		65					70			75				80		
	Leu	Asn	Leu	Asn	His	Leu	Leu	Gln	Tyr	Asn	Pro	Lys	Gln	Ile	Asp	Ile
15								85			90			95		
	Ser	Asn	Thr	Arg	Ala	Thr	Gln	Ser	Gln	Phe	Glu	Lys	Trp	Tyr	Glu	Gly
								100			105			110		
20	Val	Arg	Asp	Asp	Tyr	Gly	Leu	Asn	Asp	Asn	Glu	Met	Gln	Val	Met	Leu
							115			120			125			
	Asn	Gly	Leu	Met	Val	Trp	Cys	Ile	Glu	Asn	Gly	Thr	Ser	Pro	Asp	Ile
							130			135			140			
25	Ser	Gly	Val	Trp	Val	Met	Val	Asp	Gly	Glu	Thr	Gln	Val	Asp	Tyr	Pro
							145			150			155			160
	Ile	Lys	Pro	Leu	Ile	Glu	His	Ala	Thr	Pro	Ser	Phe	Arg	Gln	Ile	Met
30								165			170			175		
	Ala	His	Phe	Ser	Asn	Ala	Ala	Glu	Ala	Tyr	Ile	Ala	Met	Arg	Asn	Ala
								180			185			190		
35	Thr	Glu	Arg	Tyr	Met	Pro	Arg	Tyr	Gly	Ile	Lys	Arg	Asn	Leu	Thr	Asp
								195			200			205		
	Ile	Asn	Leu	Ala	Arg	Tyr	Ala	Phe	Asp	Phe	Tyr	Glu	Val	Asn	Ser	Lys
								210			215			220		

- 26 -

Xaa Pro Asp Arg Ala Arg Glu Ala His Met Gln Met Lys Alà Ala Ala
225 230 235 240

Leu Arg Asn Thr Asn Arg Arg Met Phe Gly Ile Asp Gly Ser Val Ser
5 245 250 255

Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val Asp Asp Val Asn Arg
260 265 270

10 Asp Met His Ser Leu Leu Gly Met Arg Asn
275 280

Also suitable for use in the present invention are variants of the nucleic acid molecules shown above. An example of a suitable nucleic acid is a 15 nucleic acid molecule which has a nucleotide sequence that is at least 85% similar to the nucleotide sequence of the SEQ ID NOS: 1, 3, 5, 6, 9, 11, 13, 15, 17, and 19 by basic BLAST using default parameters analysis, or which hybridizes to the nucleotide sequence of SEQ ID NOS: 1, 3, 5, 6, 9, 11, 13, 15, 17, and 19 under stringent conditions characterized by a hybridization buffer comprising 5X SSC 20 buffer at a temperature of about 42°-65°C, preferably 45°C.

Fragments of genes encoding PRSV-CP are particularly useful in the present invention. Fragments capable of use in the present invention can be produced by several means. In one method, subclones of the gene encoding the CP of choice are produced by conventional molecular genetic manipulation by 25 subcloning gene fragments. In another approach, based on knowledge of the primary structure of the protein, fragments of a PRSV-CP encoding gene may be synthesized by using the PCR technique together with specific sets of primers chosen to represent particular portions of the protein. These, then, would be cloned into an appropriate vector in either the sense or antisense orientation.

30 Another example of suitable fragments of the nucleic acids of the present invention are fragments of the genes which have been identified as conserved ("con") regions of the CP proteins, or alternatively, those portions of PRSV-CP nucleotide sequences that have been identified as variable ("var") regions. Sequences identified using DNAMega alignment program as either 35 variable or conserved in a PRSV-CP gene can be amplified using standard PCR

methods using forward and reverse primers designed to amplify the region of choice and which include a restriction enzyme sequence to allow ligation of the PCR product into a vector of choice. Combinations of amplified conserved and variable region sequences can be ligated into a single vector to create a "cassette" 5 which contains a plurality of DNA molecules in one vector. The use of conserved and variable regions of PRSV-CP DNA is further detailed below in the Examples.

The present invention also relates to a DNA construct that contains a DNA molecule encoding for a PRSV-CP isolated from any of a variety of PRSV strains, most preferably the TH, KA, KE, YK, ME, BR, JA, OA, and VE strains.

10 This involves incorporating one or more of the nucleic acid molecules of the present invention, or a suitable portion thereof, of the nucleic acid corresponding to SEQ ID NOS: 1, 3, 5, 6, 9, 11, 13, 15, 17, and 19 into host cells using conventional recombinant DNA technology. Generally, this involves inserting the nucleic acid molecule into an expression system to which the nucleic acid 15 molecule is heterologous (i.e., not normally present). The heterologous nucleic acid molecule is inserted into the expression system which includes the necessary elements for the transcription and translation of the inserted protein coding sequences.

The nucleic acid molecules of the present invention may be 20 inserted into any of the many available expression vectors and cell systems using reagents that are well known in the art. Suitable vectors include, but are not limited to, the following viral vectors such as lambda vector system gt11, gt WES.tB, Charon 4, and plasmid vectors such as pBR322, pBR325, pACYC177, pACYC1084, pUC8, pUC9, pUC18, pUC19, pLG339, pR290, pKC37, pKC101, 25 SV 40, pBluescript II SK +/- or KS +/- (see "Stratagene Cloning Systems" Catalog (1993) from Stratagene, La Jolla, CA, which is hereby incorporated by reference in its entirety), pQE, pIH821, pGEX, pET series (see Studier et. al., "Use of T7 RNA Polymerase to Direct Expression of Cloned Genes," Gene Expression Technology vol. 185 (1990), which is hereby incorporated by 30 reference in its entirety), and any derivatives thereof. Recombinant molecules can be introduced into cells via transformation, particularly transduction, conjugation, mobilization, or electroporation. The DNA sequences are cloned into the vector using standard cloning procedures in the art, as described by Sambrook et al.,

Molecular Cloning: A Laboratory Manual, Second Edition, Cold Spring Harbor Press, NY (1989), and Ausubel, F. M. et al. (1989) Current Protocols in Molecular Biology, John Wiley & Sons, New York, N.Y., which are hereby incorporated by reference in their entirety.

- 5 In preparing a DNA vector for expression, the various DNA sequences may normally be inserted or substituted into a bacterial plasmid. Any convenient plasmid may be employed, which will be characterized by having a bacterial replication system, a marker which allows for selection in a bacterium, and generally one or more unique, conveniently located restriction sites.
- 10 Numerous plasmids, referred to as transformation vectors, are available for plant transformation. The selection of a vector will depend on the preferred transformation technique and target species for transformation. A variety of vectors are available for stable transformation using *Agrobacterium tumefaciens*, a soilborne bacterium that causes crown gall. Crown gall are characterized by
- 15 tumors or galls that develop on the lower stem and main roots of the infected plant. These tumors are due to the transfer and incorporation of part of the bacterium plasmid DNA into the plant chromosomal DNA. This transfer DNA ("T-DNA") is expressed along with the normal genes of the plant cell. The plasmid DNA, pTi, or Ti-DNA, for "tumor inducing plasmid," contains the *vir*
- 20 genes necessary for movement of the T-DNA into the plant. The T-DNA carries genes that encode proteins involved in the biosynthesis of plant regulatory factors, and bacterial nutrients (opines). The T-DNA is delimited by two 25 bp imperfect direct repeat sequences called the "border sequences." By removing the oncogene and opine genes, and replacing them with a gene of interest, it is possible to
- 25 transfer foreign DNA into the plant without the formation of tumors or the multiplication of *Agrobacterium tumefaciens* (Fraley, et al., "Expression of Bacterial Genes in Plant Cells," Proc. Nat'l Acad. Sci. 80:4803-4807 (1983), which is hereby incorporated by reference in its entirety).

 Further improvement of this technique led to the development of

30 the binary vector system (Bevan, M., "Binary *Agrobacterium* Vectors for Plant Transformation," Nucleic Acids Res. 12:8711-8721 (1984), which is hereby incorporated by reference in its entirety). In this system, all the T-DNA sequences (including the borders) are removed from the pTi, and a second vector containing

- 29 -

T-DNA is introduced into *Agrobacterium tumefaciens*. This second vector has the advantage of being replicable in *E. coli* as well as *A. tumefaciens*, and contains a multiclonal site that facilitates the cloning of a transgene. An example of a commonly used vector is pBin19 (Frisch, et al., "Complete Sequence of the 5 Binary Vector Bin19," Plant Molec. Biol. 27:405-409 (1995), which is hereby incorporated by reference in its entirety). Any appropriate vectors now known or later described for genetic transformation are suitable for use in the present invention.

U.S. Patent No. 4,237,224 issued to Cohen and Boyer, which is 10 hereby incorporated by reference in its entirety, describes the production of expression systems in the form of recombinant plasmids using restriction enzyme cleavage and ligation with DNA ligase. These recombinant plasmids are then introduced by means of transformation and replicated in unicellular cultures including prokaryotic organisms and eukaryotic cells grown in tissue culture.

15 Certain "control elements" or "regulatory sequences" are also incorporated into the vector-construct. These include non-translated regions of the vector, promoters, and 5' and 3' untranslated regions which interact with host cellular proteins to carry out transcription and translation. Such elements may vary in their strength and specificity. Depending on the vector system and host 20 utilized, any number of suitable transcription and translation elements, including constitutive and inducible promoters, may be used.

25 A constitutive promoter is a promoter that directs expression of a gene throughout the development and life of an organism. Examples of some constitutive promoters that are widely used for inducing expression of transgenes include the nopaline synthase ("NOS") gene promoter, from *Agrobacterium tumefaciens*, (U.S. Patent 5034322 to Rogers et al., which is hereby incorporated by reference in its entirety), the cauliflower mosaic virus ("CaMV") 35S and 19S promoters (U.S. Patent No. 5,352,605 to Fraley et al., which is hereby incorporated by reference in its entirety), the enhanced CaMV35S promoter ("enh 30 CaMV35S"), the figwort mosaic virus full-length transcript promoter ("FMV35S"), those derived from any of the several actin genes, which are known to be expressed in most cells types (U.S. Patent No. 6,002,068 to Privalle et al., which is hereby incorporated by reference in its entirety), and the ubiquitin

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promoter ("ubi"), which is a gene product known to accumulate in many cell types.

An inducible promoter is a promoter that is capable of directly or indirectly activating transcription of one or more DNA sequences or genes in response to an inducer. In the absence of an inducer, the DNA sequences or genes will not be transcribed. The inducer can be a chemical agent, such as a metabolite, growth regulator, herbicide or phenolic compound, or a physiological stress directly imposed upon the plant such as cold, heat, salt, toxins, the action of a pathogen or disease agent such as a virus or fungus. A plant cell containing an inducible promoter may be exposed to an inducer by externally applying the inducer to the cell or plant such as by spraying, watering, heating, or by exposure to the operative pathogen. An example of an appropriate inducible promoter for use in the present invention is a glucocorticoid-inducible promoter ("GIP") (Schena et al., "A Steroid-Inducible Gene Expression System for Plant Cells," 15 Proc. Natl. Acad. Sci. 88:10421-5 (1991), which is hereby incorporated by reference in its entirety). Other useful promoters include promoters capable of expressing potyvirus proteins in an inducible manner or in a tissue-specific manner in certain cell types where infection is known to occur. These include, for example, the inducible promoters from phenylalanine ammonia lyase, chalcone 20 synthase, extensin, pathogenesis-related protein, and wound-inducible protease inhibitor from potato. Other examples of such tissue specific promoters include seed, flower, or root specific promoters as are well known in the field (U.S. Patent No. 5,750,385 to Shewmaker et al., which is hereby incorporated by reference in its entirety). For a review on maximizing gene expression, see Roberts and Lauer, 25 Methods in Enzymology 68:473 (1979), which is hereby incorporated by reference in its entirety.

The particular promoter selected is preferably capable of causing sufficient expression of the DNA coding sequences to which it is operably linked, to result in the production of amounts of the proteins effective to provide viral 30 resistance, but not so much as to be detrimental to the cell in which they are expressed. The actual choice of the promoter is not critical, as long as it has sufficient transcriptional activity to accomplish the expression of the preselected proteins, where expression is desired, and subsequent conferral of viral resistance

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to the plants. The promoters selected should be capable of functioning in tissues including, but not limited to, epidermal, vascular, and mesophyll tissues.

The nucleic acid construct of the present invention also includes an operable 3' regulatory region, which provides a functional poly(A) addition signal (AATAAA) 3' of its translation termination codon. This is selected from among those which are capable of providing correct transcription termination and polyadenylation of mRNA for expression in the host cell of choice, operably linked to a DNA molecule which encodes for a protein of choice. A number of 3' regulatory regions are known to be operable in plants. Exemplary 3' regulatory regions include, without limitation, the nopaline synthase 3' regulatory region (Fraley, et al., "Expression of Bacterial Genes in Plant Cells," Proc. Nat'l Acad. Sci. USA 80:4803-4807 (1983), which is hereby incorporated by reference in its entirety) and the cauliflower mosaic virus 3' regulatory region (Odell, et al., "Identification of DNA Sequences Required for Activity of the Cauliflower Mosaic Virus 35S Promoter," Nature 313(6005):810-812 (1985), which is hereby incorporated by reference in its entirety). Virtually any 3' regulatory region known to be operable in plants would suffice for proper expression of the coding sequence of the nucleic acid construct of the present invention.

A vector of choice, suitable promoter, and an appropriate 3' regulatory region can be ligated together to produce the expression systems which contain the nucleic acids of the present invention, or suitable fragments thereof, using well known molecular cloning techniques as described in Sambrook et al., Molecular Cloning: A Laboratory Manual, Second Edition, Cold Spring Harbor Press, NY (1989), and Ausubel et al. (1989) Current Protocols in Molecular Biology, John Wiley & Sons, New York, N.Y., which are hereby incorporated by reference in their entirety.

Once the isolated nucleic acid molecules encoding the various papaya ringspot virus coat proteins or polypeptides, as described above, have been cloned into an expression system, they are ready to be incorporated into a host cell. Such incorporation can be carried out by the various forms of transformation noted above, depending upon the vector/host cell system. Suitable host cells include, but are not limited to, bacteria, virus, yeast, mammalian cells, insect, plant, and the like.

Accordingly, another aspect of the present invention relates to a recombinant plant cell containing one or more of the PRSV-CP nucleic acids of the present invention. Basically, this method is carried out by transforming a plant cell with a nucleic acid construct of the present invention under conditions effective to yield transcription of the DNA molecule in response to the promoter.

5 Methods of transformation may result in transient or stable expression of the DNA under control of the promoter. Preferably, the nucleic acid construct of the present invention is stably inserted into the genome of the recombinant plant cell as a result of the transformation, although transient expression can serve an

10 important purpose, particularly when the plant under investigation is slow-growing.

Plant tissue suitable for transformation include without limitation, leaf tissue, root tissue, meristems, zygotic and somatic embryos, callus, protoplasts, tassels, pollen, embryos, anthers, and the like. The means of transformation chosen is that most suited to the tissue to be transformed.

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Transient expression in plant tissue is often achieved by particle bombardment (Klein et al., "High-Velocity Microprojectiles for Delivering Nucleic Acids Into Living Cells," Nature 327:70-73 (1987), which is hereby incorporated by reference in its entirety). In this method, tungsten or gold microparticles (1 to 2 μm in diameter) are coated with the DNA of interest and then bombarded at the tissue using high pressure gas. In this way, it is possible to deliver foreign DNA into the nucleus and obtain a temporal expression of the gene under the current conditions of the tissue. Biologically active particles (e.g., dried bacterial cells containing the vector and heterologous DNA) can also be propelled into plant cells (U.S. Patent Nos. 4,945,050, 5,036,006, and 5,100,792, all to Sanford et al., which are hereby incorporated by reference in their entirety). For papaya, particle gun bombardment has been a particularly successful method (Fitch, M.M., "Stable Transformation of Papaya Via Micro-Projectile Bombardment," Plant Cell Rep. 9:189 (1990), and Fitch et al., "Somatic

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An appropriate method of stably introducing the nucleic acid construct into plant cells is to infect a plant cell with *Agrobacterium tumefaciens* or *Agrobacterium rhizogenes* previously transformed with the nucleic acid construct. As described above, the Ti (or R1) plasmid of *Agrobacterium* enables 5 the highly successful transfer of a foreign DNA into plant cells. Yet another method of introduction is fusion of protoplasts with other entities, either minicells, cells, lysosomes or other fusible lipid-surfaced bodies (Fraley, et al., Proc. Natl. Acad. Sci. USA 79:1859-63 (1982), which is hereby incorporated by reference in its entirety). The DNA molecule may also be introduced into the plant cells by 10 electroporation (Fromm et al., Proc. Natl. Acad. Sci. USA 82:5824 (1985), which is hereby incorporated by reference in its entirety). In this technique, plant protoplasts are electroporated in the presence of plasmids containing the expression cassette. Electrical impulses of high field strength reversibly permeabilize biomembranes allowing the introduction of the plasmids.

15 Electroporated plant protoplasts reform the cell wall, divide, and regenerate. The precise method of transformation is not critical to the practice of the present invention. Any method that results in efficient transformation of the host cell of choice is appropriate for practicing the present invention.

After transformation, the transformed plant cells must be 20 regenerated. Plant regeneration from cultured protoplasts is described in Evans et al., Handbook of Plant Cell Cultures, Vol. 1: (MacMillan Publishing Co., New York, 1983); Vasil I.R. (ed.), Cell Culture and Somatic Cell Genetics of Plants, Acad. Press, Orlando, Vol. I, 1984, and Vol. III (1986), and Fitch et al., "Somatic Embryogenesis and Plant Regeneration from Immature Zygotic Embryos of 25 Papaya (*Carica papaya L.*)," Plant Cell Rep. 9:320 (1990), which are hereby incorporated by reference in their entirety.

It is known that practically all plants can be regenerated from 30 cultured cells or tissues, including but not limited to, all major species of sugarcane, sugar beets, cotton, fruit trees, and legumes. Means for regeneration vary from species to species of plants, but generally, a suspension of transformed protoplasts or a petri plate containing explants is first provided. Callus tissue is formed and shoots may be induced from callus and subsequently rooted. Alternatively, embryo formation can be induced

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in the callus tissue. These embryos germinate as natural embryos to form plants. The culture media will generally contain various amino acids and hormones, such as auxin and cytokinins. Efficient regeneration will depend on the medium, on the genotype, and on the history of the culture. If these three variables are controlled,
5 then regeneration is usually reproducible and repeatable.

Preferably, transformed cells are first identified using a selection marker simultaneously introduced into the host cells along with the nucleic acid construct of the present invention. Suitable selection markers include, without limitation, markers encoding for antibiotic resistance, such as the *nptII* gene which
10 confers kanamycin resistance (Fraley, et al., Proc. Natl. Acad. Sci. USA 80:4803-4807 (1983), which is hereby incorporated by reference in its entirety), and the genes which confer resistance to gentamycin, G418, hygromycin, streptomycin, spectinomycin, tetracycline, chloramphenicol, and the like. Cells or tissues are grown on a selection medium containing the appropriate antibiotic, whereby
15 generally only those transformants expressing the antibiotic resistance marker continue to grow. Other types of markers are also suitable for inclusion in the expression cassette of the present invention. For example, a gene encoding for herbicide tolerance, such as tolerance to sulfonylurea is useful, or the *dhfr* gene, which confers resistance to methotrexate (Bourouis et al., EMBO J. 2:1099-1104
20 (1983), which is hereby incorporated by reference in its entirety). Similarly, "reporter genes," which encode for enzymes providing for production of an identifiable compound are suitable. The most widely used reporter gene for gene fusion experiments has been *uidA*, a gene from *Escherichia coli* that encodes the β-glucuronidase protein, also known as GUS (Jefferson et al., "GUS Fusions: β
25 Glucuronidase as a Sensitive and Versatile Gene Fusion Marker in Higher Plants," EMBO J. 6:3901-3907 (1987), which is hereby incorporated by reference in its entirety). Similarly, enzymes providing for production of a compound identifiable by luminescence, such as luciferase, are useful. The selection marker employed will depend on the target species; for certain target species, different antibiotics,
30 herbicide, or biosynthesis selection markers are preferred.

Plant cells and tissues selected by means of an inhibitory agent or other selection marker are then tested for the acquisition of the viral gene by Southern blot hybridization analysis, using a probe specific to the viral genes

contained in the given cassette used for transformation (Sambrook et al., "Molecular Cloning: A Laboratory Manual," Cold Spring Harbor, New York: Cold Spring Harbor Press (1989), which is hereby incorporated by reference in its entirety).

- 5 The presence of a viral coat protein gene can also be detected by immunological assays, such as the double-antibody sandwich assays described by Namba et al., "Expression of the Gene Encoding the Coat Protein of Cucumber Mosaic Virus (CMV) Strain WL appears to Provide Protection to Tobacco Plants Against Infection by Several Different CMV Strains," Gene 107:181-188 (1991),
10 which is hereby incorporated by reference in its entirety, as modified by Clark et al., "Characteristics Of the Microplate Method for Enzyme-Linked Immunosorbent Assay For the Detection of plant Viruses," J. Gen. Virol. 34, 475-83 (1977), which is hereby incorporated by reference in its entirety. Potyvirus resistance can also be assayed via infectivity studies as generally described by
15 Namba et al., "Protection of Transgenic Plants Expressing the Coat Protein Gene of Watermelon Virus ii or Zucchini Yellow Mosaic Virus Against Potyviruses," Phytopath. 82:940946 (1992), which is hereby incorporated by reference in its entirety, wherein plants are scored as symptomatic when any inoculated leaf shows vein-clearing, mosaic, or necrotic symptoms.
20 After the expression cassette is stably incorporated in transgenic plants, it can be transferred to other plants by sexual crossing. Any of a number of standard breeding techniques can be used, depending upon the species to be crossed. Once transgenic plants of this type are produced, the plants themselves can be cultivated in accordance with conventional procedure so that the nucleic acid construct is present in the resulting plants. Alternatively, transgenic seeds or propagules (e.g., cuttings) are recovered from the transgenic plants. These seeds can then be planted in the soil and cultivated using conventional procedures to produce transgenic plants.
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30 The present invention also relates to DNA constructs which contain a plurality of DNA molecules which are derived from one or more genes which encode a papaya ringspot viral coat protein. The *PRSV-CP* DNA molecules may be derived from one or more strains, including, but not limited to, TH, KE, KA, ME, YK, BR, JA, OA, and VE. Some of the *PRSV-CP* DNA molecules may be a

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fragment of the nucleic acid sequence of the CP(s) of choice which by itself is too short, i.e., does not contain sufficient nucleotide sequence, to impart its respective trait when placed in a vector and used to transform plant cells as described above. Collectively, however, this plurality of DNA molecules impart their trait 5 to the transformed plant. The trait which is imparted is resistance to the PRSV strain from which any given DNA molecule in the construct is derived. Suitable nucleic acids for this construct include fragments of a PRSV CP-encoding DNA molecule, of any strain, including but not limited to, TH, KE, KA, ME, YK, BR, JA, OA, and VE. The DNA molecules are inserted in the construct as less than 10 full-length DNA, preferably in the range of about 200 bp of the full-length PRSV CP DNA molecule. The 200 bp fragments are preferably chosen from the conserved and variable regions of CP-encoding DNA. There is no need to include separate promoters for each of the fragments; only a single promoter is required. Moreover, such viral gene fragments can preferably be incorporated in a single 15 expression system to produce transgenic plants with a single transformation event.

The present invention also relates to a DNA construct containing a fusion gene which includes a trait DNA molecule which has a length insufficient to independently impart a desired trait to plants transformed with the trait molecule, operatively coupled to a silencer molecule effective to achieve post- 20 transcriptional gene silencing. The trait DNA molecule and the silencer molecule collectively impart the trait to plants transformed with the construct. The trait DNA molecules of this DNA construct are derived from a gene encoding a papaya ringspot viral coat protein from a papaya ringspot virus strains which include, but are not limited to TH, KE, KA, ME, YK, BR, JA, OA, and VE. The fragments of 25 trait DNA molecules are subcloned into the fusion gene cassette. Suitable DNA fragments are those of about 200 bp which derive from the variable and conserved regions of the CP-encoding molecules of choice. The silencer molecule of the construct of the present invention can be selected from virtually any nucleic acid which effects gene silencing. This involves the cellular mechanism to degrade 30 mRNA homologous to the transgene mRNA. The silencer DNA molecule can be heterologous to the plant, need not interact with the trait DNA molecule in the plant, and can be positioned 3' to the trait DNA molecule. For example, the silencer DNA molecule can be a viral cDNA molecule, including, without

limitation, a gene encoding a replicase, a movement protein, or a nucleocapsid protein; a green fluorescence protein encoding DNA molecule, a plant DNA molecule, or combinations thereof.

In any of the constructs of the present invention, the DNA
5 molecule conferring disease resistance can be positioned within the DNA construct in the sense (5'→3') orientation. Alternatively, it can have an antisense (3'→5') orientation. Antisense RNA technology involves the production of an RNA molecule that is complementary to the messenger RNA molecule of a target gene. The antisense RNA can potentially block all expression of the targeted
10 gene. In the anti-virus context, plants are made to express an antisense RNA molecule corresponding to a viral RNA (that is, the antisense RNA is an RNA molecule which is complementary to a "plus" (+) sense RNA species encoded by an infecting virus). Such plants may show a slightly decreased susceptibility to infection by that virus. Such a complementary RNA molecule is termed antisense
15 RNA.

It is possible for the DNA construct of the present invention to be configured so that the trait and silencer DNA molecules encode RNA molecules which are translatable. As a result, that RNA molecule will be translated at the ribosomes to produce the protein encoded by the DNA construct. Production of
20 proteins in this manner can be increased by joining the cloned gene encoding the DNA construct of interest with synthetic double-stranded oligonucleotides which represent a viral regulatory sequence (i.e., a 5' untranslated sequence) (U.S. Patent No. 4,820,639 to Gehrke, and U.S. Patent No. 5,849,527 to Wilson, which are hereby incorporated by reference in their entirety).

25 Alternatively, the DNA construct of the present invention can be configured so that the trait and silencer DNA molecules encode mRNA which is not translatable. This is achieved by introducing into the DNA molecule one or more premature stop codons, adding one or more bases (except multiples of 3 bases) to displace the reading frame, removing the translation initiation codon, etc.
30 See U.S. Patent No. 5,583,021 to Dougherty et al., which is hereby incorporated by reference in its entirety. The subject DNA construct can be incorporated in cells using conventional recombinant DNA technology, such as described in detail above.

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Another aspect of the present invention is a method to confer resistance to PRSV to plants. This involves transforming susceptible plants with one or more of the nucleic acid constructs of the present invention, testing for transformation using a marker inherent in the vector, selecting transgenics, and 5 regenerating and reproducing the transgenic plants as described above. The expression system of the present invention can be used to transform virtually any plant tissue under suitable conditions. Transformed cells can be regenerated into whole plants such that the PRSV-transgene imparts resistance to PRSV in the intact transgenic plants. In either case, the plant cells transformed with the 10 recombinant DNA expression system of the present invention are grown and caused to express the DNA molecule or molecules in the constructs of the present invention, and, thus, to impart papaya ringspot resistance.

While not wishing to be bound by theory, by use of the constructs of the present invention, it is believed that post-transcriptional gene silencing is 15 achieved. More particularly, the silencer DNA molecule is believed to boost the level of heterologous RNA within the cell above a threshold level. This activates the degradation mechanism by which viral resistance is achieved.

Transgenic plants which show post-transcription gene silencing-derived resistance establish the highly resistant state and prevent virus replication. 20 A chimeric transgene consisting of a silencer DNA (e.g., *GFP*) fused with various small nontranslatable fragment viral genome would be preferred for viral resistance. There are several advantages. First, the silencer DNA can increase the induced gene silencing. Second, the chimeric nature of the gene would provide multiple virus resistance. Third, nontranslatable construction produces no protein, 25 thus reducing the possible complementation of naturally occurring mutants and transencapsidation of other viruses. Fourth, the small fragment also reduces the possibility of recombination with other viral genomes.

Absent a complete understanding of the mechanism(s) of viral resistance conferred through this type of genetic manipulation, optimization of the 30 production of viral resistant transgenics is still under study. Thus, the degree of resistance imparted to a given transgenic plant (high, medium, or low efficacy) is unpredictable. However, it has been noted that when combinations of viral gene expression cassettes are placed in the same binary plasmid, and that multigene

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cassette containing plasmid is transformed into a plant, the viral genes all exhibit substantially the same degrees of efficacy when present in transgenic plants. For example, if one examines numerous transgenic lines containing two different intact viral gene cassettes, the transgenic line will be immune to infection by both 5 viruses. Likewise if a transgenic line exhibits a delay in symptom development to one virus, it will also exhibit a delay in symptom development to the second virus. Finally, if a transgenic line is susceptible to one of the viruses it will be susceptible to the other. This phenomenon is unexpected. If there were not a correlation between the efficacy of each gene in these multiple gene constructs, 10 this approach as a tool in plant breeding would probably be prohibitively difficult to use. The probability of finding a line with useful levels of expression can range from 10-50%, depending on the species involved (U.S. Patent No. 6,002,072 to McMaster et al., which is hereby incorporated by reference in its entirety).

15 The present invention will be further described by reference to the following detailed examples.

EXAMPLES

Example 1– Amplification and Cloning of CP Variable Region DNAs

20 Total RNA was extracted from PRSV-infected papaya plants. Different *PRSV-CP* gene fragments, each about 200 bp, from Taiwan (YK), Keaau (KE), and Thailand (TH) strains were amplified by reverse-transcription and polymerase-chain-reaction (RT-PCR) and extracted from agarose gels. The primers used to amplify the variable region of the PRSV-CP gene of strains YK, 25 KE, and TH are shown in Table 1.

- 40 -

Table 1

PRSV Strain	Product (bp)	Primer position	Primer Sequence (SEQ ID NO)
YKvar	209	21-39	5' GAGAtctaga <u>TAATGATAACCGGTCTGAATGAGAAG</u> 3' (SEQ ID NO: 21)
			3' YkvarXba
KEvar	209	212-229	5' GGATctcgag <u>AGATCATCTTATCAGTAA</u> 3' (SEQ ID NO: 22)
			3' KEvarXho
THvar	206	21-39	5' TAGActcgag <u>TGCTGGTTGAATGAAAAAA</u> 3' (SEQ ID NO: 23)
			3' KEvarSma
		211-229	5' CGATccggg <u>GAATCAACTTATCAGTAA</u> 3' (SEQ ID NO: 24)
			3' THvarSma
		209-226	5' TATAccggg <u>TGCTGGTCTTAATGAGAAG</u> 3' (SEQ ID NO: 25)
			3' THvarBam

5 Restriction enzyme sequence is shown in small letters; the stop codon is shown in caps, without underline; viral sequences are underlined.

Following amplification using conventional PCR techniques, the amplified fragments were digested with the appropriate restriction enzymes. A restriction enzyme *Xba*I-*Xho*I digested YK fragment (209 bp) was first ligated 10 into the pEPJ vector. A *Xho*I-*Sma*I digested KE fragment (209 bp) was ligated behind (i.e., at the 3' end of) the YK fragment and then a *Sma*I-*Bam*HI digested TH fragment (206 bp) was ligated behind the KE. The resultant clone, pEPJ-YKT, shown in Figure 1A, contains the variable region of CP from YK-KE-TH in the 5'→ 3' direction. Following a *Hind*III-*Kpn*I restriction digest, the pEPJ-YKT 15 expression cassette was ligated into the *Hind*III-*Kpn*I cloning site of transformation vector pGA482G, shown in Figure 1B, resulting in clone pTi-EPJ-YKT. Cesium chloride purified pTi-EPJ-YKT was then used for host cell transformation by particle gun bombardment.

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Example 2 -Cloning of CP Variable Regions into Silencer Construct

Fragments *Xba*I/*Bam*HI from pEPJ-YKT were ligated into other expression vectors pNP, shown in Figure 2A, and pGFP, shown in Figure 2B, creating pNP-YKT and pGFP-YKT, respectively. "M1/2 NP" shown in Figure 5 2A refers to a fragment consisting of approximately one half (387-453 bp) of the gene encoding the nucleocapsid protein ("N" or "NP" gene) of the viral genome of the tomato spotted wilt virus ("TSWV"), a tospovirus that causes crop damage worldwide. Expression of large fragments (approximately 1/2 or greater) of the N gene of TSWV have been shown to confer high levels of resistance to TSWV-BL 10 in 20-51% of R1 plants transformed with the fragment, and tolerance to tospovirus infection in 4-22% of R1 plants isolate but not to the distantly related Impatiens necrotic spot virus ("INSV") (Law et al., "The M RNA of Impatiens Necrotic Spot Tospovirus (Bunyaviridae) Has an Ambisense Genomic Organization," *Virology*, 188:732-41 (1992), which is hereby incorporated by reference in its entirety) or 15 groundnut ringspot virus ("GRSV") (Pang et al., "The Biological Properties of a Distinct Tospovirus and Sequence Analysis of Its mRNA," *Phytopathology*, 83:728-33 (1993), which is hereby incorporated by reference in its entirety). The N gene of TSWV is an example of a gene derived from the viral genome that is useful as a silencer molecule in the nucleic acid constructs of the present 20 invention. Restriction enzyme *Hind*III/*Kpn*I digested fragments from these two expression vectors were then ligated into the *Hind*III/*Kpn*I cloning site of the transformation vector pGA482G, resulting in clones pTi-NP-YKT and pTi-GFP-YKT. Cesium chloride purified pTi-NP-YKT and pTi-GFP-YKT were then used for host cell transformation by particle gun bombardment.

25

Example 3 -Amplification and Cloning of CP Conserved Region DNAs

Total RNA was extracted from PRSV-infected papaya plants. Different *PRSV-CP* gene fragments, each about 200 bp, from Keaau (KE) and Thailand (TH) were amplified by RT-PCR. The primers used to amplify the 30 conserved region of the PRSV-CP gene of strains KE and TH are shown in Table 2.

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Table 2

PRSV Strain	Product (bp)	Primer position	Primer Sequence (SEQ ID NO)
KEcon	203	649-686	5'TCAAtctagagtcgac <u>GCTAGATATGCTTCGAC</u> 3' (SEQ ID NO: 27)
		834-851	5'AAGTctcgaggcgac <u>CCCAGGAGAGAGTGCATG</u> 3' (SEQ ID NO: 28)
THcon	203	646-683	5'AATAccgggg <u>GCTAGATATGCTTCGAC</u> 3' (SEQ ID NO: 29)
		831-848	5'TTATggatcc <u>CCTAGGAGAGAGTGCATG</u> 3 (SEQ ID NO: 30)

Restriction enzyme sequence is shown in small letters; the stop codon is shown in caps, without underline; viral sequences are underlined.

5

Constructs containing the silencer molecule 1/2 NP are shown in Figures 3A-G. These constructs are designated herein as clone pNP-X_n, where "X" denotes of PRSV strain from which the CP DNA is derived, and "n" represents the number fragments of "X" in the cassette. When the DNA is inserted in the sense orientation, "X" is the first initial of the strain, for example, "K" for KE, "T" for TH. When a fragment is inserted in the antisense orientation, the strain acronym is flipped, for example, KE becomes EK, and "X" becomes the first initial of the antisense designation. For example, for an antisense fragment of KE, "X" becomes "E." Translatable and nontranslatable forms of the DNA molecule are further designated with the prefix "TL" and "NTL", respectively.

10 Clone pNP-K, shown in Figure 3A, was obtained by ligating a single 203 bp *Xba*I/*Xho*I digested KE DNA fragment in a sense orientation into the expression vector pNP containing the 365 bp M1/2NP DNA molecule. Clone pNP-KK, shown in Figure 3B, and pNP-EE, shown Figure 3C, containing sense and antisense KE fragments, respectively, were obtained by ligating a *Sall*

20 digested KE DNA fragment into pNP-K. Clone pNP-KKTC, shown in Figure 3D, pNP-KKTV, shown in Figure 3E, pNP-EETC, shown in Figure 3F; and pNP-EETV, shown in Figure 3G, were obtained by ligating a *Sma*I/*Bam*HI digested KE

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fragment from the conserved region (KEcon) or from the variable region (KEvar) into pNP-KK or pNP-EE.

The pNP clones were *Hind*III /*Kpn*I digested from the expression vectors, and ligated into the *Hind*III/*Kpn*I cloning site of the transformation vector 5 pGA482G, resulting in clones pTi-NP-K, pTi-NP-KK, pTi-NP-EE, pTi-NP-KKTC, pTi-NP-KKTV, pTi-NP-EETC and pTi-NP-EETV. Cesium chloride purified pTi-NP-clones were then used for host cell transformation by particle gun bombardment.

10 **Example 4 - Amplification and Cloning of Full Length Translatable and Nontranslatable KE**

Two full-length KE-CP constructs, shown in Figure 4, start from the first CP cut site which is 60 nt upstream from the second CP cut site. The 15 primers used for amplification and construction of pEPJ-TL KE and pEPJ-NTL KE are shown in Table 3.

Table 3

PRSV Strain	Product (bp)	Primer Sequence (SEQ ID NO)
TL KE	921	5'AGCTAAccatgg <u>AATCAAGGAGCACTGATGATTATC</u> 3' (SEQ ID NO: 31)
		5'ATTTggatcccg <u>ggGTTGCGCATGCCAGGAGAGAG</u> 3' (SEQ ID NO: 32)
NTL KE	921	5' AGCTAAccatgg <u>AATAATGGAGCACTGATGATTATC</u> 3' (SEQ ID NO: 33)
		5'ATTTggatcccg <u>ggGTTGCGCATGCCAGGAGAGAG</u> 3' (SEQ ID NO: 34)

20 Restriction enzyme sequence is shown in small letters; the stop codon is shown in caps, without underline; viral sequences are underlined.

Following amplification, the *Nco*I/*Bam*HI digested PCR KEC^P fragments were ligated into pEPJ vector, as shown in Figure 4. Using

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*Hind*II/*Kpn*I, the expression cassette was then subcloned into the transformation vector pGA482G.

Example 5 -Amplification and Cloning of MEX CP

5 The primers used for amplification and preparation of construct pEPJ-MEX CP are shown in Table 4.

Table 4

PRSV Strain	Product (bp)	Primer Sequence (SEQ ID NO)
NTL Mex	855	5'CGAtctagaccatggAATAATGAT <u>CCAAGAAATGAAGC</u> 3' (SEQ ID NO: 35)
		5'CTTAggatcc <u>GTTGCGCATACCCAGGAGAGA</u> 3' 3' (SEQ ID NO: 36)

10 Restriction enzyme sequence is shown in small letters; the stop codon is shown in caps, without underline; viral sequences are underlined.

Example 6 - Transformation of Papaya with PRSV-CP DNA Constructs

Papaya embryos were bombarded with DNA constructs prepared as
 15 described above and shown in Figures 2-5. The transformation procedure was followed as described in Cai et al., "A Protocol for Efficient Transformation and Regeneration of *Carica papaya L. In Vitro*," *Cell Devel. Biol-Plant* 35: 61-69 (1999), which is hereby incorporated by reference in its entirety. Plasmid DNA was purified by ethidium bromide CsCl gradient (Ausubel et al., "CsCl/Ethidium
 20 Bromide Preparations of Plasmid DNA," *Current Protocols in Molec Biol.* unit 2.9.1-2.9.20 (1995), which is hereby incorporated by reference in its entirety), ethanol precipitated and suspended in water. Immature zygotic embryos were extracted from seeds of immature green 'Sunrise' or 'Kapoho' papaya and placed on induction medium and kept in the dark. Zygotic embryos with their somatic
 25 embryo clusters were placed on Whatman #2 filter paper and spread. The somatic embryos were allowed to proliferate, and following this, the embryos were spread firmly onto fresh filter paper and bombarded with tungsten-coated plasmid DNA. Seven days after bombardment, materials were transferred to induction medium

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containing kanamycin at 75 mg/L. After four weeks, the kanamycin level was raised to 150 mg/L. After a few weeks in kanamycin medium, actively growing embryo clusters were transferred to kanamycin-free medium. When the embryos developed a pale ivory color and appeared as finger-like extensions, they were
5 transferred to maturation medium for two to four weeks. Mature somatic embryos were transferred to germination medium and then developed into plantlets with dark green leaves and root initials. Those plantlets were transferred to baby jars with rooting medium and transferred to the greenhouse.

Transgenic lines from the germination medium were analyzed by
10 PCR to confirm that the virus gene was in the plantlets. Northern blots were carried out to detect the level of RNA expressed in transgenic lines, and the copy number of the transgene in the transgenic plants was determined by Southern blot analysis.

Following transfer to the greenhouse, transgenic plants were
15 challenged with the KE strain of PRSV. Plants were thereafter monitored for viral symptoms. If no disease symptoms appeared after approximately 4 weeks post-inoculation, those plants were challenged with a different PRSV strain to test for cross-resistance.

20 **Example 7 - Resistance Imparted to PRSV by Transgenes**

219 transgenic lines containing the various PRSV DNA constructs of the present invention, as described above, were transferred to the greenhouse. Inoculation with KE virus was carried out on 90 plant lines transformed with at least one KE-containing DNA construct. Of those 90 lines challenged with PRSV-
25 KE, 26 lines showed resistance and 64 lines were susceptible.

Although preferred embodiments have been depicted and described in detail herein, it will be apparent to those skilled in the relevant art that various modifications, additions, substitutions, and the like can be made without departing from the spirit of the invention and these are therefore considered to be within the
30 scope of the invention as defined in the claims which follow.

WHAT IS CLAIMED:

1. An isolated nucleic acid molecule encoding a papaya ringspot virus coat protein, wherein the nucleic acid molecule either: 1) has a
5 nucleic acid sequence of SEQ ID NO: 1; 2) encodes an amino acid having SEQ ID NO: 2; 3) has a nucleotide sequence that is at least 85% similar to the nucleotide sequence of SEQ ID NO: 1 by basic BLAST using default parameters analysis; or 4) hybridizes to the nucleotide sequence of SEQ ID NO: 1 under stringent conditions characterized by a hybridization buffer comprising 5X SSC buffer at a
10 temperature of 45°C.
2. A DNA construct comprising:
the nucleic acid molecule according to claim 1 and
an operably linked promoter and 3' regulatory region.
15
3. A DNA expression vector comprising:
the DNA construct according to claim 2.
4. A host cell transduced with a DNA construct according to
20 claim 2.
5. A host cell according to claim 4, wherein the cell is selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a plant cell.
25
6. A transgenic plant transformed with a DNA construct according to claim 2.
7. A transgenic plant according to claim 6, wherein the plant is papaya.
30
8. A transgenic plant seed transformed with a DNA construct according to claim 2.

9. A transgenic plant seed according to claim 8, wherein the plant is papaya.

5 10. An isolated nucleic acid molecule encoding a papaya ringspot virus coat protein wherein the nucleic acid molecule either: 1) has a nucleic acid sequence of SEQ ID NO: 3; 2) encodes an amino acid having SEQ ID NO: 4; 3) has a nucleotide sequence that is at least 85% similar to the nucleotide sequence of SEQ ID NO: 3 by basic BLAST using default parameters analysis; or
10 4) hybridizes to the nucleotide sequence of SEQ ID NO: 3 under stringent conditions characterized by a hybridization buffer comprising 5X SSC buffer at a temperature of 45°C.

15 11. A DNA construct comprising:
the nucleic acid molecule according to claim 10 and
an operably linked promoter and 3' regulatory region.

20 12. An expression vector comprising the DNA construct of
claim 11.

25 13. A host cell transduced with a DNA construct according to
claim 11.

14. A host cell according to claim 13, wherein the cell is selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a plant cell.

30 15. A transgenic plant transformed with a DNA construct according to claim 11.

16. A transgenic plant according to claim 15, wherein the plant is papaya.

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17. A transgenic plant seed transformed with a DNA construct according to claim 11.

5 18. A transgenic plant seed according to claim 17, wherein the plant is papaya.

10 19. An isolated nucleic acid molecule encoding a papaya ringspot virus coat protein, wherein the nucleic acid molecule either: 1) has a nucleic acid sequence of SEQ ID NO: 5; 2) encodes an amino acid having SEQ ID NO: 7; 3) has a nucleotide sequence that is at least 85% similar to the nucleotide sequence of SEQ ID NO: 5 by basic BLAST using default parameters analysis; or 4) hybridizes to the nucleotide sequence of SEQ ID NO: 5 under stringent conditions characterized by a hybridization buffer comprising 5X SSC buffer at a 15 temperature of 45°C.

20 20. A DNA construct comprising:
the nucleic acid molecule according to claim 19 and
an operably linked promoter and 3' regulatory region.

21. A DNA expression vector comprising:
the DNA construct according to claim 20.

22. A host cell transduced with a DNA construct according to 25 claim 20.

23. A host cell according to claim 22, wherein the cell is selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a plant cell.

30 24. A transgenic plant transformed with a DNA construct according to claim 20.

25. A transgenic plant according to claim 24, wherein the plant
is papaya.

5 26. A transgenic plant seed transformed with a DNA construct
according to claim 20.

27. A transgenic plant seed according to claim 26, wherein the
plant is papaya.

10 28. An isolated nucleic acid molecule encoding a papaya
ringspot virus coat protein wherein the nucleic acid molecule either: 1) has a
nucleic acid sequence of SEQ ID NO: 6; 2) encodes an amino acid having SEQ ID
NO: 8; 3) has a nucleotide sequence that is at least 85% similar to the nucleotide
15 sequence of SEQ ID NO: 6 by basic BLAST using default parameters analysis; or
4) hybridizes to the nucleotide sequence of SEQ ID NO: 6 under stringent
conditions characterized by a hybridization buffer comprising 5X SSC buffer at a
temperature of 45°C.

20 29. A DNA construct comprising:
the nucleic acid molecule according to claim 28 and
an operably linked promoter and 3' regulatory region.

30. An expression vector comprising the DNA construct of
25 claim 29.

31. A host cell transduced with a DNA construct according to
claim 29.

30 32. A host cell according to claim 31, wherein the cell is
selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a
plant cell.

- 50 -

33. A transgenic plant transformed with a DNA construct according to claim 29.

34. A transgenic plant according to claim 33, wherein the plant
5 is papaya.

35. A transgenic plant seed transformed with a DNA construct according to claim 29.

10 36. A transgenic plant seed according to claim 35, wherein the plant is papaya.

37. An isolated nucleic acid molecule encoding a papaya ringspot virus coat protein, wherein the nucleic acid molecule either: 1) has a
15 nucleic acid sequence of SEQ ID NO: 11; 2) encodes an amino acid having SEQ ID NO: 12; 3) has a nucleotide sequence that is at least 85% similar to the nucleotide sequence of SEQ ID NO: 11 by basic BLAST using default parameters analysis; or 4) hybridizes to the nucleotide sequence of SEQ ID NO: 11 under stringent conditions characterized by a hybridization buffer comprising 5X SSC
20 buffer at a temperature of 45°C.

25 38. A DNA construct comprising:
the nucleic acid molecule according to claim 37 and
an operably linked promoter and 3' regulatory region.

39. A DNA expression vector comprising:
the DNA construct according to claim 38.

40. A host cell transduced with a DNA construct according to
30 claim 38.

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41. A host cell according to claim 40, wherein the cell is selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a plant cell.

5 42. A transgenic plant transformed with a DNA construct according to claim 38.

43. A transgenic plant according to claim 42, wherein the plant is papaya.

10 44. A transgenic plant seed transformed with a DNA construct according to claim 38.

45. A transgenic plant seed according to claim 44, wherein the 15 plant is papaya.

46. An isolated nucleic acid molecule encoding a papaya ringspot virus coat protein wherein the nucleic acid molecule either: 1) has a nucleic acid sequence of SEQ ID NO: 13; 2) encodes an amino acid having SEQ 20 ID NO: 14; 3) has a nucleotide sequence that is at least 85% similar to the nucleotide sequence of SEQ ID NO: 13 by basic BLAST using default parameters analysis; or 4) hybridizes to the nucleotide sequence of SEQ ID NO: 13 under stringent conditions characterized by a hybridization buffer comprising 5X SSC buffer at a temperature of 45°C.

25 47. A DNA construct comprising:
the nucleic acid molecule according to claim 46 and
an operably linked promoter and 3' regulatory region.

30 48. An expression vector comprising the DNA construct of
claim 47.

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49. A host cell transduced with a DNA construct according to
claim 47.

50. A host cell according to claim 49, wherein the cell is
selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a
plant cell.

51. A transgenic plant transformed with a DNA construct
according to claim 47.

10 52. A transgenic plant according to claim 51, wherein the plant
is papaya.

15 53. A transgenic plant seed transformed with a DNA construct
according to claim 47.

54. A transgenic plant seed according to claim 53, wherein the
plant is papaya.

20 55. An isolated nucleic acid molecule encoding a papaya
ringspot virus coat protein, wherein the nucleic acid molecule either: 1) has a
nucleic acid sequence of SEQ ID NO: 15; 2) encodes an amino acid having SEQ
ID NO: 16; 3) has a nucleotide sequence that is at least 85% similar to the
nucleotide sequence of SEQ ID NO: 15 by basic BLAST using default parameters
analysis; or 4) hybridizes to the nucleotide sequence of SEQ ID NO: 15 under
stringent conditions characterized by a hybridization buffer comprising 5X SSC
buffer at a temperature of 45°C.

30 56. A DNA construct comprising:
the nucleic acid molecule according to claim 55 and
an operably linked promoter and 3' regulatory region.

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57. A DNA expression vector comprising:
the DNA construct according to claim 56.

58. A host cell transduced with a DNA construct according to
5 claim 56.

59. A host cell according to claim 58, wherein the cell is
selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a
plant cell.

10

60. A transgenic plant transformed with a DNA construct
according to claim 56.

15 61. A transgenic plant according to claim 60, wherein the plant
is papaya.

62. A transgenic plant seed transformed with a DNA construct
according to claim 56.

20 63. A transgenic plant seed according to claim 62, wherein the
plant is papaya.

25 64. An isolated nucleic acid molecule encoding a papaya
ringspot virus coat protein wherein the nucleic acid molecule either: 1) has a
nucleic acid sequence of SEQ ID NO: 17; 2) encodes an amino acid having SEQ
ID NO: 18; 3) has a nucleotide sequence that is at least 85% similar to the
nucleotide sequence of SEQ ID NO: 17 by basic BLAST using default parameters
analysis; or 4) hybridizes to the nucleotide sequence of SEQ ID NO: 17 under
stringent conditions characterized by a hybridization buffer comprising 5X SSC
30 buffer at a temperature of 45°C.

65. A DNA construct comprising:

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the nucleic acid molecule according to claim 64 and
an operably linked promoter and 3' regulatory region.

5 66. An expression vector comprising the DNA construct of
claim 65.

67. A host cell transduced with a DNA construct according to
claim 65.

10 68. A host cell according to claim 67, wherein the cell is
selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a
plant cell.

15 69. A transgenic plant transformed with a DNA construct
according to claim 65.

70. A transgenic plant according to claim 69, wherein the plant
is papaya.

20 71. A transgenic plant seed transformed with a DNA construct
according to claim 65.

72. A transgenic plant seed according to claim 71, wherein the
plant is papaya.

25 73. An isolated nucleic acid molecule encoding a papaya
ringspot virus coat protein wherein the nucleic acid molecule either: 1) has a
nucleic acid sequence of SEQ ID NO: 19; 2) encodes an amino acid having SEQ
ID NO: 20; 3) has a nucleotide sequence that is at least 85% similar to the
30 nucleotide sequence of SEQ ID NO: 19 by basic BLAST using default parameters
analysis; or 4) hybridizes to the nucleotide sequence of SEQ ID NO: 19 under

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stringent conditions characterized by a hybridization buffer comprising 5X SSC buffer at a temperature of 45°C.

5 74. A DNA construct comprising:
the nucleic acid molecule according to claim 73 and
an operably linked promoter and 3' regulatory region.

10 75. An expression vector comprising the DNA construct of
claim 74.

15 76. A host cell transduced with a DNA construct according to
claim 74.

20 77. A host cell according to claim 76, wherein the cell is
selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a
plant cell.

25 78. A transgenic plant transformed with a DNA construct
according to claim 74.

30 79. A transgenic plant according to claim 78, wherein the plant
is papaya.

25 80. A transgenic plant seed transformed with a DNA construct
according to claim 74.

30 81. A transgenic plant seed according to claim 80, wherein the
plant is papaya.

35 82. A DNA construct comprising:
a plurality of trait DNA molecules at least some of which have a
length that is insufficient to impart that trait to plants transformed with that trait

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DNA molecule, but said plurality of trait DNA molecules collectively impart their traits to plants transformed with said DNA construct and effect silencing of the DNA construct, wherein the trait is disease resistance and the trait DNA molecules are derived from a gene encoding a papaya ringspot virus coat protein
5 in a papaya ringspot virus strain selected from the group consisting of TH, KE, KA, ME, YK, BR, JA, OA, VE, and PA.

83. A DNA construct according to claim 82, wherein one or more of the trait DNA molecules are selected from the group consisting of the
10 variable regions and conserved regions of said papaya ringspot viral coat proteins.

84. The DNA construct according to claim 82, wherein one or more of the trait DNA molecules are in the sense (5'→3') orientation.

15 85. The DNA construct according to claim 82, wherein one or more of the trait DNA molecules are inserted in the antisense (3'→5') orientation.

86. An expression vector comprising:
the DNA construct according to claim 82.

20 87. A host cell transduced with a DNA construct according to
claim 82.

88. A host cell according to claim 87, wherein the cell is
25 selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a plant cell.

89. A transgenic plant transformed with a DNA construct
according to claim 82.

30 90. A transgenic plant according to claim 89, wherein the plant
is papaya.

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91. A transgenic plant seed transformed with a DNA construct according to claim 82.

92. A transgenic plant seed according to claim 91, wherein the
5 plant is papaya.

93. A DNA construct comprising:
a fusion gene comprising:
a trait DNA molecule which has a length that is insufficient to
10 independently impart a desired trait to plants transformed with said trait DNA
molecule and
a silencer DNA molecule effective to achieve post-transcriptional
gene silencing and operatively coupled to said trait DNA molecule, wherein said
trait DNA molecule and said silencer DNA molecule collectively impart the trait
15 to the plants transformed with said DNA construct, and wherein the trait DNA
molecules are derived from a gene encoding a papaya ringspot viral coat protein
from a papaya ringspot virus strain selected from the group consisting of TH, KE,
KA, ME, YK, BR, JA, OA, VE, and PA.

20 94. A DNA construct according to claim 93, further
comprising:

a promoter sequence operatively coupled to said fusion gene and
a termination sequence operatively coupled to said fusion gene to
end transcription.

25 95. A DNA construct according to claim 93, wherein said
silencer DNA molecule is selected from the group consisting of a viral DNA
molecule, a fluorescence protein encoding DNA molecule, a plant DNA molecule,
a viral gene silencer, and combinations thereof.

30 96. An expression vector comprising:
the DNA construct according to claim 93.

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97. A host cell transduced with a DNA construct according to
claim 93.

98. A host cell according to claim 97, wherein the cell is
5 selected from the group consisting of a bacterial cell, a virus, a yeast cell, and a
plant cell.

99. A transgenic plant transformed with a DNA construct
according to claim 93.

10 100. A transgenic plant according to claim 36, wherein the plant
is papaya.

101. A transgenic plant seed transformed with a DNA construct
15 according to claim 93.

102. A transgenic plant according to claim 101, wherein the
plant is papaya.

20 103. A method of imparting resistance to papaya plants against
papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 2.

25 104. A method of imparting resistance to papaya plants against
papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 11.

30 105. A method of imparting resistance to papaya plants against
papaya ringspot virus comprising:

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transforming a papaya plant with a DNA construct
according to claim 20.

106. A method of imparting viral resistance to papaya plants
5 against papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 29.

107. A method of imparting viral resistance to papaya plants
10 against papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 38.

108. A method of imparting viral resistance to papaya plants
15 against papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 47.

109. A method of imparting viral resistance to papaya plants
20 against papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 56.

110. A method of imparting viral resistance to papaya plants
25 against papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 65.

111. A method of imparting viral resistance to papaya plants
30 against papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 74.

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112. A method of imparting viral resistance to papaya plants against papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 82.

5

113. A method of imparting viral resistance to papaya plants against papaya ringspot virus comprising:
transforming a papaya plant with a DNA construct according to
claim 93.

10

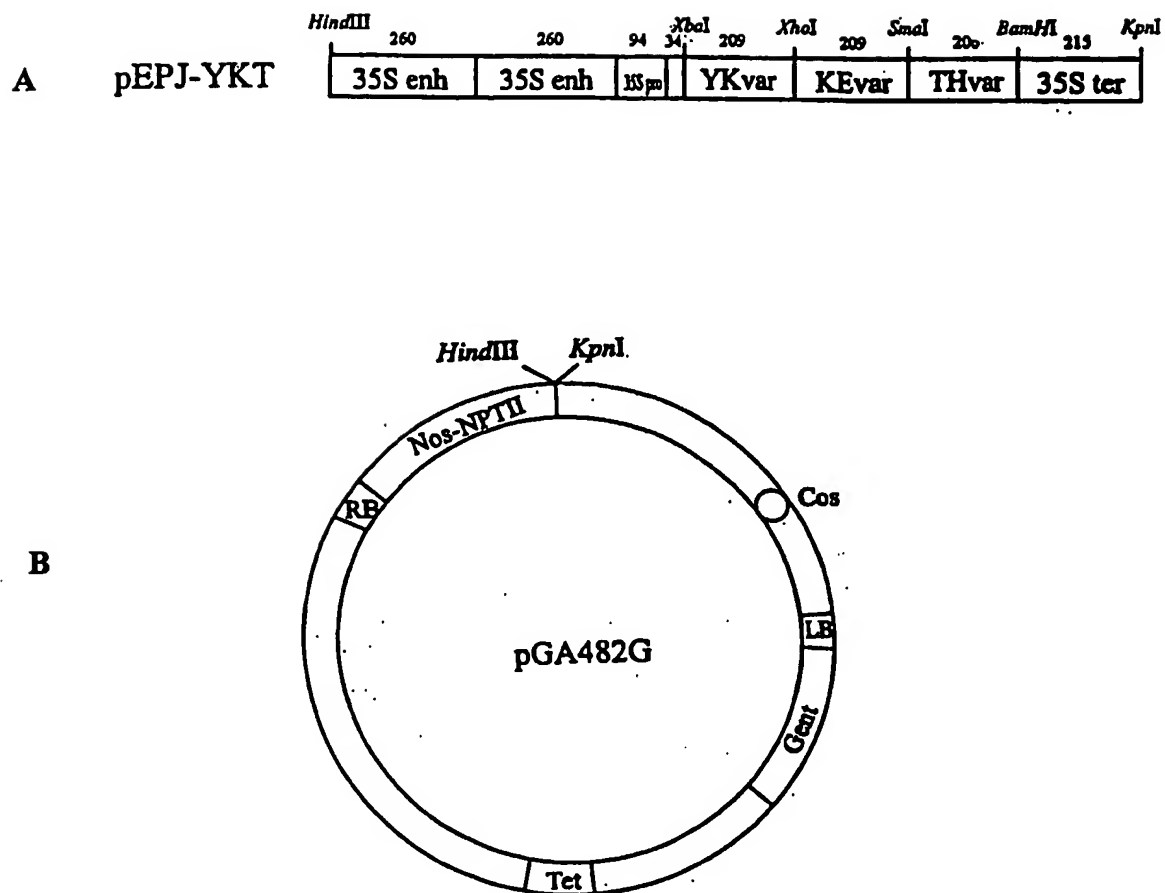


FIGURE 1

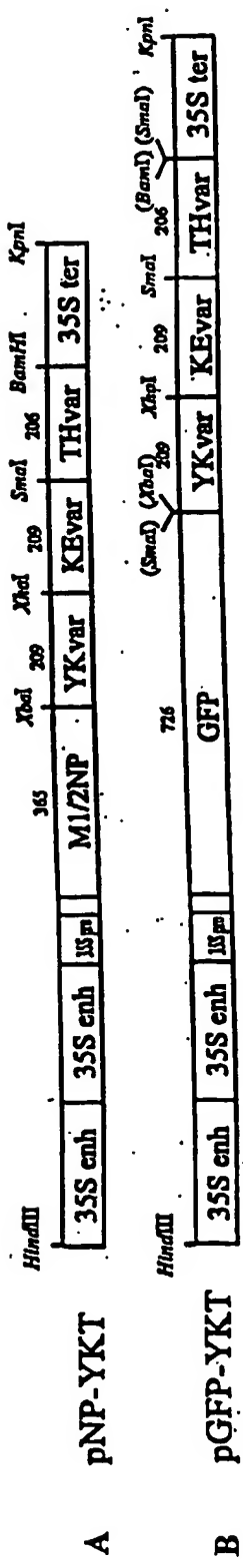
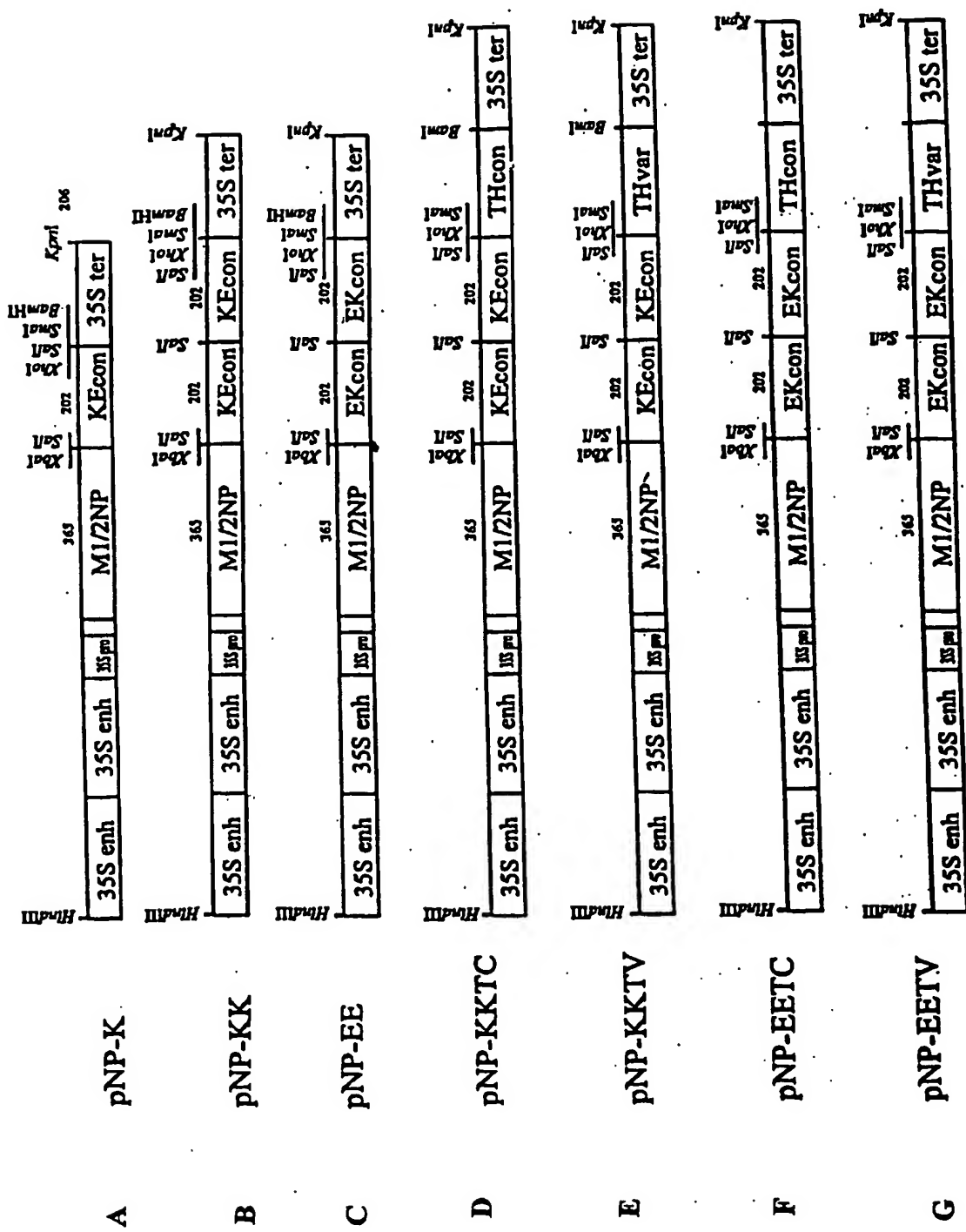


FIGURE 2

**FIGURE 3**

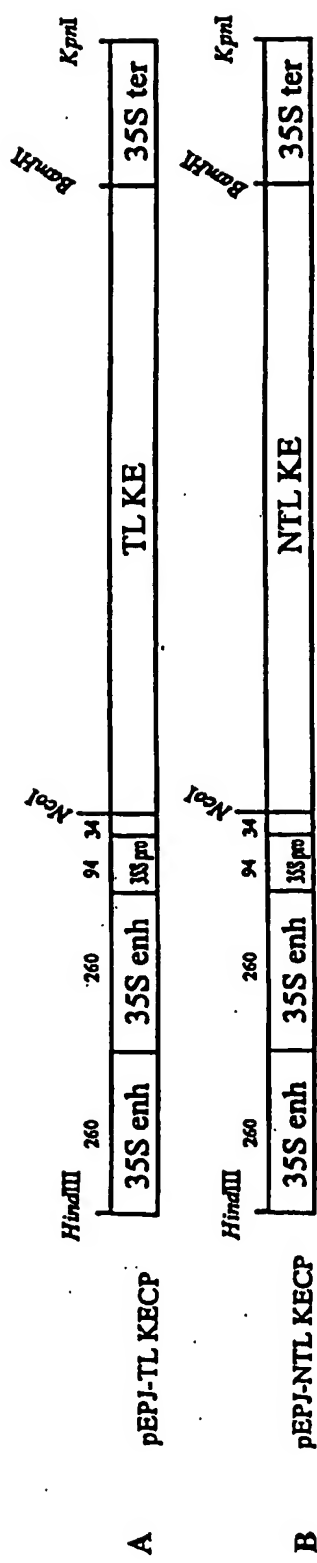
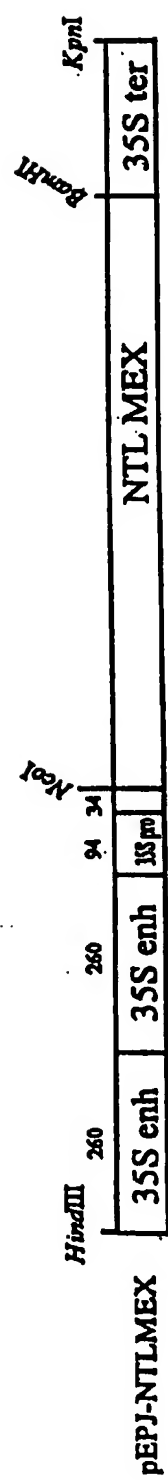


FIGURE 4

**FIGURE 5**

SEQUENCE LISTING

<110> Gonsalves, Dennis
Chiang, Chu-Hui
Tennant, Paula F.
Gonsalves, Carol V.
Sarindu, Nonglak
Souza, Jr., Manoel Teixeira
Nickel, Osmar
Munoz, Gustavo Alberto Fermin
Saxena, Sanjay
Cai, Wengi

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35 40 45

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180 185 190

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Tyr Gly Ile Lys Arg Asn Leu Thr Asp Val Ser Leu Ala Arg Tyr Ala
 225 230 235 240

Phe Asp Phe Tyr Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu
 245 250 255

Ala His Met Gln Met Lys Ala Ala Ala Leu Arg Asn Thr Ser Arg Arg
 260 265 270

Met Phe Gly Met Asp Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu
 275 280 285

Arg His Thr Val Glu Asp Val Asn Arg Asp Met His Ser Leu Leu Gly
 290 295 300

Met Arg Asn
 305

<210> 8
<211> 287
<212> PRT
<213> PRSV-KE-CP2

<400> 8
Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Leu Lys Glu
 1 5 10 15

Lys Glu Lys Gln Lys Glu Lys Glu Lys Glu Lys Gln Lys Glu Lys Gly
 20 25 30

Lys Asp Asp Ala Ser Asp Glu Asn Asp Val Ser Thr Ser Thr Lys Thr
 35 40 45

Gly Glu Arg Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Ala
 50 55 60

Val Pro Arg Ile Lys Ser Phe Thr Asp Lys Leu Ile Leu Pro Arg Ile
 65 70 75 80

Lys Gly Lys Thr Val Leu Asn Leu Ser His Leu Leu Gln Tyr Asn Pro
 85 90 95

Gln Gln Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu
 100 105 110

Lys Trp Tyr Glu Gly Val Arg Asp Asp Tyr Gly Leu Asn Asp Asn Glu
 115 120 125

Met Gln Val Met Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly
 130 135 140

Thr Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr
 145 150 155 160

Gln Val Asp Tyr Pro Thr Lys Pro Leu Ile Glu His Ala Thr Pro Ser
 165 170 175

Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile
 180 185 190

Ala Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys
 195 200 205

Arg Asn Leu Thr Asp Val Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr
 210 215 220

Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln
 225 230 235 240

Met Lys Ala Ala Ala Leu Arg Asn Thr Ser Arg Arg Met Phe Gly Met
 245 250 255

Asp Gly Ser Val Ser Asn Lys Glu Asn Thr Glu Arg His Thr Val
 260 265 270

Glu Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn
 275 280 285

<210> 9
<211> 864
<212> DNA
<213> PRSV-YK-CP

<400> 9
tctaaaaatg aagctgtgga taccggctcg aatgagaagc tcaaagaaaa agaaaagcag 60
aaagaaaaag aaaaagataa acaacaagat aaagacaatg atggagctag tgacggaaac 120
gatgtgtcaa ctagcacaaa aactggagag agagataggg atgtcaatgc cggaactagt 180
ggaaccttca ctgttccgag gataaagtca tttactgata agatgatctt accaagaatt 240
aaggaaaaaa ctgtccttaa tttaaatcat cttcttcagt ataatccgaa acaagttgac 300
atctcaaaca ctcgcgccac tcaatctcaa tttgagaagt ggtatgaggg aqtgagaaat 360
gattatggcc ttaatgataa cgaaatgcaa gtaatgttaa atggtttgat gggttgggt 420
atcgaaaatg gtacatctcc agatatatct ggtgtctggg ttatgatgga tggggaaacc 480

caagtcgatt atccccattaa acctttgatt gaacacgcaa ctccttcatt taggcaaatc 540
 atggctcaact ctagtaacgc ggcagaggca tacatcgca agaggaatgc aactgagaag 600
 tacatgccgc ggtatgaaat caagagaaat ttgactgaca ttagtctcgc tagatatgct 660
 ttcgatttct atgaggtgaa ttcgaaaaca cctgataggg ctcgtgaagc tcataatgcag 720
 atgaaggctg cagcgctacg caatactaatt cgcaaaatgt ttgaaatgga cggcagtgtc 780
 agtaacaagg aagaaaacac ggagagacac acagtgaaag atgtcaacag agacatgcac 840
 tctctcctgg gtatgcgcaa ttga 864

<210> 10
<211> 287
<212> PRT
<213> PRSV-YK-CP

<400> 10

Ser	Lys	Asn	Glu	Ala	Val	Asp	Thr	Gly	Leu	Asn	Glu	Lys	Leu	Lys	Glu
1															
															15

Lys	Glu	Lys	Gln	Lys	Glu	Lys	Glu	Lys	Asp	Lys	Gln	Gln	Asp	Lys	Asp
20															
															30

Asn	Asp	Gly	Ala	Ser	Asp	Gly	Asn	Asp	Val	Ser	Thr	Ser	Thr	Lys	Thr
35															
															45

Gly	Glu	Arg	Asp	Arg	Asp	Val	Asn	Ala	Gly	Thr	Ser	Gly	Thr	Phe	Thr
50															
															60

Val	Pro	Arg	Ile	Lys	Ser	Phe	Thr	Asp	Lys	Met	Ile	Leu	Pro	Arg	Ile
65															
															80

Lys	Gly	Lys	Thr	Val	Leu	Asn	Leu	Asn	His	Leu	Leu	Gln	Tyr	Asn	Pro
85															
															95

Lys	Gln	Val	Asp	Ile	Ser	Asn	Thr	Arg	Ala	Thr	Gln	Ser	Gln	Phe	Glu
100															
															110

Lys	Trp	Tyr	Glu	Gly	Val	Arg	Asn	Asp	Tyr	Gly	Leu	Asn	Asp	Asn	Glu
115															
															125

Met	Gln	Val	Met	Leu	Asn	Gly	Leu	Met	Val	Trp	Cys	Ile	Glu	Asn	Gly
130															
															140

Thr	Ser	Pro	Asp	Ile	Ser	Gly	Val	Trp	Val	Met	Met	Asp	Gly	Glu	Thr
145															
															160

Gln	Val	Asp	Tyr	Pro	Ile	Lys	Pro	Leu	Ile	Glu	His	Ala	Thr	Pro	Ser
165															
															175

Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile
 180 185 190

Ala Lys Arg Asn Ala Thr Glu Lys Tyr Met Pro Arg Tyr Gly Ile Lys
 195 200 205

Arg Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr
 210 215 220

Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln
 225 230 235 240

Met Lys Ala Ala Ala Leu Arg Asn Thr Asn Arg Lys Met Phe Gly Met
 245 250 255

Asp Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val
 260 265 270

Glu Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn
 275 280 285

<210> 11
<211> 855
<212> DNA
<213> PRSV-ME-CP

<400> 11

tccaaagaatg aagctgtgga tgctggtttgc aatgaaaaac tcaaaagaaaa agaaaaacag 60
 aaagaaaaag aaaaacaaaa agaaaaagaa aaagacaatg ctatgtacgg aaatgtatgt 120
 tcgacttagca caaaaactgg agagaaagat agagatgtca atgtcggaac tagtggaaact 180
 ttcaactgttc cgagaattaa atcatttact gataagatga ttctaccgag aattaaggga 240
 aagactgtcc ttaatttaaa tcatcttctt cagttataatc cgcaacaaat tgatatttct 300
 aacactcgtg ccactcagtc acaatttgag aaatggatg agggagtgag gaatgattat 360
 ggtctgaatg ataatgaaat gcaagtgtatg ctgaatggct tgatggtttgc gtgtatcgag 420
 aatggtacat ctccagacat atctgggttttgc tggatggggaa aattcaagtt 480
 gactatccaa tcaaggctct aattgagcat gctacccgt catttaggca gattatggct 540
 cacttttagta acgcggcaga agcatatatt gcaaagagaa atgccactga gaggtacatg 600
 ccgcggatg gaatcaagag aaatttgact gacattagcc tcgcttagtgc cgctttcgat 660
 ttctatgagg ttaattcgaa aacacctgtatggcgtcgaa aagctcacat gcagatgaaa 720
 gctgcagcgc tgcgaaacac tagtcgcaga atgtttggta tgggcggcag tgtagtaac 780
 aaggaagaaaa acacggaaag acacacagtgc gaagatgtca atagagacat gcactctctc 840
 ctgggtatgc gcaac 855

<210> 12
<211> 285
<212> PRT

<213> PRSV-ME-CP

<400> 12

Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Leu Lys Glu
1 5 10 15

Lys Glu Lys Gln Lys Glu Lys Glu Lys Gln Lys Glu Lys Glu Lys Asp
20 25 30

Asn Ala Ser Asp Gly Asn Asp Val Ser Thr Ser Thr Lys Thr Gly Glu
35 40 45

Lys Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Thr Val Pro
50 55 60

Arg Ile Lys Ser Phe Thr Asp Lys Met Ile Leu Pro Arg Ile Lys Gly
65 70 75 80

Lys Thr Val Leu Asn Leu Asn His Leu Leu Gln Tyr Asn Pro Gln Gln
85 90 95

Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu Lys Trp
100 105 110

Tyr Glu Gly Val Arg Asn Asp Tyr Gly Leu Asn Asp Asn Glu Met Gln
115 120 125

Val Met Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly Thr Ser
130 135 140

Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Ile Gln Val
145 150 155 160

Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser Phe Arg
165 170 175

Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile Ala Lys
180 185 190

Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys Arg Asn
195 200 205

Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr Glu Val
210 215 220

Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln Met Lys
225 230 235 240

Ala Ala Ala Leu Arg Asn Thr Ser Arg Arg Met Phe Gly Met Gly Gly
 245 250 255

Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val Glu Asp
 260 265 270

Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn
 275 280 285

<210> 13

<211> 861

<212> DNA

<213> PRSV-BR-CP

<400> 13

tccaaaaatg aagctgtgga tgctggtttgc aatgaaaagc gtaaagaaca agagaaaacaa 60
 gaagaaaaag aagaaaaaca aaaaaaagaaa gaaaaagacg atgcttagtta cgaaaaacgat 120
 gtgtcaacta gcacaagaac tggagagaga gacagagatg tcaatgttgg gaccagtgg 180
 actttcactg ttccgagaac aaaatcattt actgataaga tgattttacc tagaattaag 240
 ggaaaaactg tccttaattt aaatcatctg attcagtata atccgcaaca aattgacatt 300
 tctaacaactc gtgctactca atcacaattt gagaagtggt acgaggaggatg gaggaatgtat 360
 tatggcctta atgataatga gatgcaaata gtgctaaatg gtttgatggt ttgggtgtatc 420
 gaaaacggta catctccaga catatctggt gtctgggtta tgatggatgg ggaaacccag 480
 gttgactatc caatcaagcc tttaattttag catgctactc cgtcgttttag gcaaatttatg 540
 gtcatttca gtaacgcggc agaagcatac attacaaga gaaatgtac tgagaggtac 600
 atgcccgggt atggatcaa gagaattttg actgacatta gtctgtctatc atatgctttc 660
 gatttctatg aggtgaatttgc gaaaacacccat gatagggttc gcgaaatgtca catgcagatg 720
 aaagctgcag cgctgcggaaa cactaatcgc agaatgttgc gtatggacgg cagtgttagt 780
 aacaaggaag aaaacacccggaa gagacacaca gtggaaatgt tcaatagaga catgcactct 840
 ctccctgggtta tgctactca 861

<210> 14

<211> 286

<212> PRT

<213> PRSV-BR-CP

<400> 14

Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Arg Lys Glu
 1 5 10 15

Gln Glu Lys Gln Glu Glu Lys Glu Glu Lys Gln Lys Lys Lys Glu Lys
 20 25 30

Asp Asp Ala Ser Tyr Gly Asn Asp Val Ser Thr Ser Thr Arg Thr Gly
 35 40 45

Glu Arg Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Thr Val
 50 55 60

Pro Arg Thr Lys Ser Phe Thr Asp Lys Met Ile Leu Pro Arg Ile Lys
 65 70 75 80

Gly Lys Thr Val Leu Asn Leu Asn His Leu Ile Gln Tyr Asn Pro Gln
 85 90 95

Gln Ile Asp Ile Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu Lys
 100 105 110

Trp Tyr Glu Gly Val Arg Asn Asp Tyr Gly Leu Asn Asp Asn Glu Met
 115 120 125

Gln Ile Val Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly Thr
 130 135 140

Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr Gln
 145 150 155 160

Val Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser Phe
 165 170 175

Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile Thr
 180 185 190

Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys Arg
 195 200 205

Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr Glu
 210 215 220

Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln Met
 225 230 235 240

Lys Ala Ala Ala Leu Arg Asn Thr Asn Arg Arg Met Phe Gly Met Asp
 245 250 255

Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val Glu
 260 265 270

Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn
 275 280 285

<210> 15

<211> 864

<212> DNA
 <213> PRSV-JA-CP

<400> 15
 tctaaaaatg aagctgtgga tgctgggaaa aatgaaaaagc tcaaaagaaaa agaaaaacag 60
 aaagataaaag aaaaagaaaa acaaaaaagat aaagaaaaag gagatgctag tgacggaaat 120
 gatggttcga cttagcacaaa aactggagag agagatagag atgtcaatgt tgggaccagt 180
 ggaacttcca ctgttccgag aattaaatca ttcaactgata agatggttct accaagaatt 240
 aaggaaaaaa ctgtccttaa tttaaatcat cttcttcagt ataatccaca acaaattgac 300
 atttctaaca ctcgtgccac tcagtcacaa ttgagaagt ggtacgaagg agtgaggagt 360
 gattatggcc taaatgatag tgaaatgcaa gtgacgctaa atggcttcat ggtttgggtgt 420
 atcgagaatg gtacatctcc agacatatct ggtgtctggg ttatgatgga tggggaaacc 480
 caagttgatt atccaatcaa gcctttaatt ggcacgcta ccccatcatt taggcagatt 540
 atggctcact tcagtaacgc ggcagaagca tacactgcaa agagaaatgc tactgagagg 600
 tacatgccgc ggtatgaaat caagagaaaat ttgactgaca ttagtctcgc tagatacgt 660
 ttcgatttct atgaggtgaa ttcaagaca cctgataggg ctcgtgaagc tcacatgcag 720
 atgaaagctg cagcgcgtcg aaacactaat cgccagaatgt ttgttatgga cggcagtgtt 780
 agtaacaatg aagaaaacac ggagagacac acagtggaaat atgtctatat agacatgcac 840
 tctctcctgc gtttgcgcaa ctga 864

<210> 16
 <211> 287
 <212> PRT
 <213> PRSV-JA-CP

<400> 16

Ser	Lys	Asn	Glu	Ala	Val	Asp	Ala	Gly	Leu	Asn	Glu	Leu	Lys	Glu
1														
														15

Lys	Glu	Lys	Gln	Lys	Asp	Lys	Glu	Lys	Glu	Lys	Gln	Lys	Asp	Lys	Glu
20														30	

Lys	Gly	Asp	Ala	Ser	Asp	Gly	Asn	Asp	Gly	Ser	Thr	Ser	Thr	Lys	Thr
35														45	

Gly	Glu	Arg	Asp	Arg	Asp	Val	Asn	Val	Gly	Thr	Ser	Gly	Thr	Ser	Thr
50														60	

Val	Pro	Arg	Ile	Lys	Ser	Phe	Thr	Asp	Lys	Met	Val	Leu	Pro	Arg	Ile
65											75			80	

Lys	Gly	Lys	Thr	Val	Leu	Asn	Leu	Asn	His	Leu	Leu	Gln	Tyr	Asn	Pro
85														95	

Gln	Gln	Ile	Asp	Ile	Ser	Asn	Thr	Arg	Ala	Thr	Gln	Ser	Gln	Phe	Glu
100											105			110	

Lys Trp Tyr Glu Gly Val Arg Ser Asp Tyr Gly Leu Asn Asp Ser Glu
 115 120 125

Met Gln Val Thr Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly
 130 135 140

Thr Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr
 145 150 155 160

Gln Val Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser
 165 170 175

Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Thr
 180 185 190

Ala Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys
 195 200 205

Arg Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr
 210 215 220

Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln
 225 230 235 240

Met Lys Ala Ala Ala Leu Arg Asn Thr Asn Arg Arg Met Phe Gly Met
 245 250 255

Asp Gly Ser Val Ser Asn Asn Glu Glu Asn Thr Glu Arg His Thr Val
 260 265 270

Glu Asp Val Tyr Ile Asp Met His Ser Leu Leu Arg Leu Arg Asn
 275 280 285

<210> 17
<211> 864
<212> DNA
<213> PRSV-OA-CP

<400> 17
tccaagaatg aagctgtgga tgctggttt aatgaaaaat tcaaagagaa ggaaaaacag 60
aaagaaaaag aaaaagaaaa acaaaaagag aaagaaaaag atggtgctag tgacgaaaat 120
gatgtgtcaa ctagcacaaa aactggagag agagatagag atgtcaatgt cgggaccagt 180
ggaactttca cagttccgag aattaaatca tttactgata agatgattct accgagaatt 240
aaggggaaagg ctgtccttaa tttaaatcat cttcttcagt acaatccgca acaaatcgac 300
atttctaaca ctcgtgccgc tcattcacaa tttgaaaagt ggtatgaggg agtgaggaat 360
gattatgccc ttaatgataa tgaaaatgcaa gtgatgctaa atggtttgat ggtttgggt 420
atcgagaatg gtacatctcc agacatatct ggtgtctggg taatgatgga tggggaaacc 480

caagtcgatt atccaatcaa gccttgatt gagcatgcta ctccgtcatt taggcaaatt 540
 atggctcaact ttagtaacgc ggcagaagca tacattgcga agagaaatgc tactgagagg 600
 tacatgccgc ggtatgaaat caagagaaat ttgactgaca ttagcctcgc tagatacgct 660
 ttgcactttt atgaggtgaa ttcgaaaaca cctgatacag ctcgcgaagc tcacatgcag 720
 atgaaggctg cagcgctgca aaacaccagt cgccagaatgt ttggtatgga cggcagtgtt 780
 agtaacaagg aagaaaacac ggagagacac acagtggaaat atgtcaatag agacatgcac 840
 tctctcctgg gtatgcgcaa ctaa 864

<210> 18
 <211> 287
 <212> PRT
 <213> PRSV-OA-CP

<400> 18

Ser Lys Asn Glu Ala Val Asp Ala Gly Leu Asn Glu Lys Phe Lys Glu
 1 5 10 15

Lys Glu Lys Gln Lys Glu Lys Glu Lys Glu Lys Gln Lys Glu Lys Glu
 20 25 30

Lys Asp Gly Ala Ser Asp Glu Asn Asp Val Ser Thr Ser Thr Lys Thr
 35 40 45

Gly Glu Arg Asp Arg Asp Val Asn Val Gly Thr Ser Gly Thr Phe Thr
 50 55 60

Val Pro Arg Ile Lys Ser Phe Thr Asp Lys Met Ile Leu Pro Arg Ile
 65 70 75 80

Lys Gly Lys Ala Val Leu Asn Leu Asn His Leu Leu Gln Tyr Asn Pro
 85 90 95

Gln Gln Ile Asp Ile Ser Asn Thr Arg Ala Ala His Ser Gln Phe Glu
 100 105 110

Lys Trp Tyr Glu Gly Val Arg Asn Asp Tyr Ala Leu Asn Asp Asn Glu
 115 120 125

Met Gln Val Met Leu Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly
 130 135 140

Thr Ser Pro Asp Ile Ser Gly Val Trp Val Met Met Asp Gly Glu Thr
 145 150 155 160

Gln Val Asp Tyr Pro Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser
 165 170 175

Phe Arg Gln Ile Met Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile
 180 185 190

Ala Lys Arg Asn Ala Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys
 195 200 205

Arg Asn Leu Thr Asp Ile Ser Leu Ala Arg Tyr Ala Phe Asp Phe Tyr
 210 215 220

Glu Val Asn Ser Lys Thr Pro Asp Arg Ala Arg Glu Ala His Met Gln
 225 230 235 240

Met Lys Ala Ala Ala Leu Arg Asn Thr Ser Arg Arg Met Phe Gly Met
 245 250 255

Asp Gly Ser Val Ser Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val
 260 265 270

Glu Asp Val Asn Arg Asp Met His Ser Leu Leu Gly Met Arg Asn
 275 280 285

<210> 19

<211> 885

<212> DNA

<213> PRSV-VE-CP

<220>

<221> unsure

<222> (678)

<223> M at position 678 in this sequence is either a or
 c

<400> 19

atggctgtgg atgctggttt gaatgggaag ctcaaagaaa aagagaaaaa agaaaaaagaa 60
 aaagaaaaac agaaagagaa agagaaagat gatgctagtg acgaaatga ttgtcaact 120
 agcacaaaaa ctggagagag agatagagat gtcaatattt ggaccagtgg aactttcact 180
 gtcccttagga ttaaatcatt tactgataag atgattttac cgagaattaa gggaaagact 240
 gtccttaatt taaatcatct tcttcgttat aatccgaaac aaatttgacat ttctaatact 300
 cgtgccactc agtcgaatt tgagaaatgg tatgaggggat tgagggatga ttatggcctt 360
 aatgataatg aaatgcattt gatgctaaat ggcttgcatttttttgatgcat tgagaatgg 420
 acatctccag acatatctgg tggttgggtt atggtggatg gggaaaccca agttgattat 480
 ccaatcaagc ctttaatttga gcatgctaca ccgtcatttta ggcaaatttggctcatttt 540
 agtaacgcgg cagaagcata cattgcgtatg agaaatgcta ctgagaggta catgccgcgg 600
 tatggaatca agagaaattt gactgacatc aacctagctc gatacgcttt tgatttctat 660
 gaggtgaatt cggaaacmcc tgataggct cgtgaagctc acatgcagat gaaggctgca 720
 gctttgcgaa acactaatcg cagaatgttt ggtatcgacg gcagtgttag caacaaggaa 780
 gaaaacacgg agagacacac agtggatgat gtcaatagag acatgcactc ttcctgggt 840

atgcgcaact aaatactcgc acttgtgtgt ttgtcgagcc tgact

885

<210> 20
<211> 282
<212> PRT
<213> PRSV-VE-CP

<220>
<221> UNSURE
<222> (225)
<223> Xaa at position 225 in this sequence is any amino acid

<400> 20
Met Ala Val Asp Ala Gly Leu Asn Gly Lys Leu Lys Glu Lys Glu Lys
1 5 10 15

Lys Glu Lys Glu Lys Glu Lys Gln Lys Glu Lys Glu Lys Asp Asp Ala
20 25 30

Ser Asp Gly Asn Asp Val Ser Thr Ser Thr Lys Thr Gly Glu Arg Asp
35 40 45

Arg Asp Val Asn Ile Thr Ser Gly Thr Phe Thr Val Pro Arg Ile Lys
50 55 60

Ser Phe Thr Asp Lys Met Ile Leu Pro Arg Ile Lys Gly Lys Thr Val
65 70 75 80

Leu Asn Leu Asn His Leu Leu Gln Tyr Asn Pro Lys Gln Ile Asp Ile
85 90 95

Ser Asn Thr Arg Ala Thr Gln Ser Gln Phe Glu Lys Trp Tyr Glu Gly
100 105 110

Val Arg Asp Asp Tyr Gly Leu Asn Asp Asn Glu Met Gln Val Met Leu
115 120 125

Asn Gly Leu Met Val Trp Cys Ile Glu Asn Gly Thr Ser Pro Asp Ile
130 135 140

Ser Gly Val Trp Val Met Val Asp Gly Glu Thr Gln Val Asp Tyr Pro
145 150 155 160

Ile Lys Pro Leu Ile Glu His Ala Thr Pro Ser Phe Arg Gln Ile Met
165 170 175

Ala His Phe Ser Asn Ala Ala Glu Ala Tyr Ile Ala Met Arg Asn Ala
180 185 190

Thr Glu Arg Tyr Met Pro Arg Tyr Gly Ile Lys Arg Asn Leu Thr Asp
195 200 205

Ile Asn Leu Ala Arg Tyr Ala Phe Asp Phe Tyr Glu Val Asn Ser Lys
210 215 220

Xaa Pro Asp Arg Ala Arg Glu Ala His Met Gln Met Lys Ala Ala Ala
225 230 235 240

Leu Arg Asn Thr Asn Arg Arg Met Phe Gly Ile Asp Gly Ser Val Ser
245 250 255

Asn Lys Glu Glu Asn Thr Glu Arg His Thr Val Asp Asp Val Asn Arg
260 265 270

Asp Met His Ser Leu Leu Gly Met Arg Asn
275 280

<210> 21

<211> 35

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Amplification
Oligos

<400> 21

gagatctaga taatgatacc ggtctgaatg agaag 35

<210> 22

<211> 28

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Amplification
Oligos

<400> 22

ggatctcgag agatcatctt atcagtaa 28

<210> 23
<211> 29
<212> DNA
<213> Artificial Sequence

<220>
<223> Description of Artificial Sequence: Amplification
Oligos

<400> 23
tagactcgag tgctggtttg aatgaaaaaa 29

<210> 24
<211> 28
<212> DNA
<213> Artificial Sequence

<220>
<223> Description of Artificial Sequence: Amplification
Oligos

<400> 24
cgatcccggg gaatcaactt atcagtaa 28

<210> 25
<211> 29
<212> DNA
<213> Artificial Sequence

<220>
<223> Description of Artificial Sequence: Amplification
Oligos

<400> 25
tatacccggg tgctggtctt aatgagaag 29

<210> 26
<211> 28
<212> DNA
<213> Artificial Sequence

<220>
<223> Description of Artificial Sequence: Amplification
Oligos

<400> 26

ctacggatcc aaatcatctt gtcggtaa

28

<210> 27

<211> 34

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Amplification
Oligos

<400> 27

tcaatctaga gtcgacgcta gatatgcttt cgac

34

<210> 28

<211> 34

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Amplification
Oligos

<400> 28

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